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(54) Title: TRANSFERRIN RECEPTOR GENES OF MORAXELLA

(57) Abstract

Purified and isolated nucleic acid molecules are provided which encode transferrin receptor proteins of *Moraxella*, such as *M. catarrhalis* or a fragment or an analog of the transferrin receptor protein. The nucleic acid sequence may be used to produce recombinant transferrin receptor proteins Tbp1 and Tbp2 of the strain of *Moraxella* free of other proteins of the *Moraxella* strain for purposes of diagnostics and medical treatment. Furthermore, the nucleic acid molecule may be used in the diagnosis of infection.

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TITLE OF INVENTION TRANSFERRIN RECEPTOR GENES OF MORAXELLA

5 FIELD OF INVENTION

The present invention relates to the molecular cloning of genes encoding transferrin receptor (TfR) proteins and, in particular, to the cloning of transferrin receptor genes from Moraxella (Branhamella) catarrhalis.

REFERENCE TO RELATED APPLICATION

This application is a continuation-in-part of copending United States Patent Application No. 08/778,570 filed January 3, 1997, which itself is a continuation-in-part of United States Patent Application No. 08/613,009 filed March 8, 1996.

BACKGROUND OF THE INVENTION

20 Moraxella (Branhamella) catarrhalis bacteria are Gram-negative diplococcal pathogens which are carried asymptomatically in the healthy human respiratory tract. In recent years, M. catarrhalis has been recognized as important causative agent of otitis media. addition, M. catarrhalis has been associated with 25 sinusitis, conjunctivitis, and urogenital infections, as well as with a number of inflammatory diseases of the respiratory tract in children and including pneumonia, chronic bronchitis, tracheitis, and 30 emphysema (refs. 1 to 8). (Throughout this application, various references are cited in parentheses to describe more fully the state of the art to which this invention pertains. Full bibliographic information citation is found at the end of the specification, 35 immediately preceding the claims. The disclosures of these references are hereby incorporated by reference

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into the present disclosure). Occasionally, *M. catarrhalis* invades to cause septicaemia, arthritis, endocarditis, and meningitis (refs. 9 to 13).

Otitis media is one of the most common illnesses of early childhood and approximately 80% of all children suffer at least one middle ear infection before the age of three (ref. 14). Chronic otitis media has been associated with auditory and speech impairment children, and in some cases, has been associated with learning disabilities. Conventional treatments for otitis media include antibiotic administration surgical procedures, including tonsillectomies, adenoidectomies, and tympanocentesis. In the United States, treatment costs for otitis media are estimated to be between one and two billion dollars per year.

In otitis media cases, M. catarrhalis commonly is co-isolated from middle ear fluid along Streptococcus pneumoniae and non-typable Haemophilus influenzae, which are believed to be responsible for 50% and 30% of otitis media infections, respectively. catarrhalis is believed to be responsible approximately 20% of otitis media infections (ref. 15). Epidemiological reports indicate that the number of cases of otitis media attributable to M. catarrhalis is increasing, along with the number of antibioticresistant isolates of M. catarrhalis. Thus, prior to 1970, no β -lactamase-producing M. catarrhalis isolates had been reported, but since the mid-seventies, increasing number of β -lactamase-expressing have been detected. Recent surveys suggest that 75% of clinical isolates produce β -lactamase (ref. 16, 26).

Iron is an essential nutrient for the growth of many bacteria. Several bacterial species, including M. catarrhalis, obtain iron from the host by using transferrin receptor proteins to capture transferrin. A number of bacteria including Neisseria meningitidis

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(ref. 17), N. gonorrhoeae (ref. 18), Haemophilus influenzae (ref. 19), as well as M. catarrhalis (ref. 20), produce outer membrane proteins which specifically bind human transferrin. The expression of these proteins is regulated by the amount of iron in the environment.

The two transferrin receptor proteins of *M. catarrhalis*, designated transferrin binding protein 1 (Tbp1) and transferrin binding protein 2 (Tbp2), have molecular weights of 115 kDa (Tbp1) and approximately 80 to 90 kDa (Tbp2). Unlike the transferrin receptor proteins of other bacteria which have an affinity for apotransferrin, the *M. catarrhalis* Tbp2 receptors have a preferred affinity for iron-saturated (i.e., ferri-) transferrin (ref. 21).

Μ. catarrhalis infection may lead to disease. It would be advantageous to provide a recombinant source of transferrin binding proteins as antigens in immunogenic preparations including vaccines, carriers for other antigens and immunogens and the generation of diagnostic reagents. The genes encoding transferrin binding proteins and fragments thereof are particularly desirable and useful in the identification and diagnosis of Moraxella and immunization against disease caused by M. catarrhalis and for the generation of diagnostic reagents.

SUMMARY OF THE INVENTION

The present invention is directed towards the 30 provision purified and isolated nucleic of molecules encoding a transferrin receptor of a strain of Moraxella or a fragment or an analog of the transferrin receptor protein. The nucleic acid molecules provided herein are useful for the specific detection of strains 35 Moraxella and for diagnosis of infection The purified and isolated nucleic acid Moraxella.

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molecules provided herein, such as DNA, are also useful for expressing the tbp genes by recombinant DNA means for providing, in an economical manner, purified and isolated transferrin receptor proteins as subunits, fragments or analogs thereof. The transferrin receptor, subunits or fragments thereof or thereof, as well as nucleic acid molecules encoding the same and vectors containing such nucleic acid molecules, are useful in immunogenic compositions for vaccinating against diseases caused by Moraxella, the diagnosis of infection by Moraxella and as tools for the generation of immunological reagents. Monoclonal antibodies or mono-specific antisera (antibodies) raised against the transferrin receptor protein, produced in accordance with aspects of the present invention, are useful for the diagnosis of infection by Moraxella, the specific detection of Moraxella (in, for example, in vitro and in vivo assays) and for the treatment of diseases caused by Moraxella.

In accordance with one aspect of the present invention, there is provided a purified and isolated nucleic acid molecule encoding a transferrin receptor protein of a strain of *Moraxella*, more particularly, a strain of *M. catarrhalis*, specifically *M. catarrhalis* strain 4223, Q8 or R1, or a fragment or an analog of the transferrin receptor protein.

In one preferred embodiment of the invention, the nucleic acid molecule may encode only the Tbpl protein of the Moraxella strain or only the Tbp2 protein of the Moraxella strain. In another preferred embodiment of the invention, the nucleic acid may encode a fragment of the transferrin receptor protein of a strain of Moraxella having an amino acid sequence which is conserved.

In another aspect of the present invention, there is provided a purified and isolated nucleic acid

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molecule having a DNA sequence selected from the group consisting of (a) a DNA sequence as set out in Figure 5, 6, 10, 11 or 27 (SEQ ID Nos: 1, 2, 3, 4, 5, 6, 7, 8, 45 or 46) or the complementary DNA sequence thereto; (b) a DNA sequence encoding an amino acid sequence as set out in Figure 5, 6, 10, 11 or 27 (SEQ ID Nos: 9, 10, 11, 13, 14, 15, 16 or 47) or the complementary DNA thereto; sequence and (C) a DNA sequence hybridizes under stringent conditions to any one of the DNA sequences defined in (a) or (b). The DNA sequence in (c) preferably has at least about sequence identity with any one of the DNA sequences defined in (a) and (b). The DNA sequence defined in (c) may be that encoding the equivalent transferrin receptor protein from another strain of Moraxella.

In an additional aspect, the present invention includes a vector adapted for transformation of a host, comprising a nucleic acid molecule as provided herein and may have the characteristics of a nucleotide sequence contained within vectors LEM3-24, pLEM3, pLEM25, pLEM23, SLRD-A, DS-1698-1-1, DS-1754-1, pSLRD2, pSLRD3, pSLRD4 and pSLRD5.

The vector may be adapted for expression of the encoded transferrin receptor, fragments or analogs thereof, in a heterologous or homologous host, in either lipidated or non-lipidated form. Accordingly, further aspect of the present invention provides an expression vector adapted for transformation of a host comprising a nucleic acid molecule as provided herein and expression means operatively coupled to the nucleic acid molecule for expression by the host transferrin receptor protein or the fragment or analog of the transferrin receptor protein. In embodiments of this aspect of the invention, the nucleic molecule may . encode substantially transferrin receptor protein, only the Tbpl protein,

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only the Tbp2 protein of the Moraxella strain fragments of the Tbpl or Tbp2 proteins. The expression means may include a promoter and a nucleic acid portion encoding a leader sequence for secretion from the host of the transferrin receptor protein or the fragment or the analog of the transferrin receptor protein. expression means also may include a nucleic acid portion encoding a lipidation signal for expression from the host of a lipidated form of the transferrin receptor protein or the fragment or the analog of the transferrin receptor protein. The host may be selected from, for example, Escherichia coli. Bordetella, Bacillus. Haemophilus, Moraxella, fungi, yeast or baculovirus and Semliki Forest virus expression systems may be used. particular embodiment, the plasmid adapted for expression of Tbpl is pLEM29 and that for expression of is pLEM33. Further vectors include SLRD35-A and SLRD-35-B.

In an additional aspect of the invention, there is provided a transformed host containing an expression vector as provided herein. The invention further includes a recombinant transferrin receptor protein or fragment or analog thereof of a strain of *Moraxella* producible by the transformed host.

Such recombinant transferrin receptor protein may be provided in substantially pure form according to a further aspect of the invention, which provides a method of forming a substantially pure recombinant transferrin receptor protein, which comprises growing transformed host provided herein to express transferrin receptor protein inclusion as bodies. purifying the inclusion bodies free from material and soluble proteins, solubilizing transferrin receptor protein from the purified inclusion bodies, and purifying the transferrin receptor protein free from other solubilized materials. The substantially pure

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recombinant transferrin receptor protein may comprise Tbpl alone, Tbp2 alone or a mixture thereof. The recombinant protein is generally at least about 70% pure, preferably at least about 90% pure.

Further aspects of the present invention, therefore, provide recombinantly-produced Tbpl protein of a strain of Moraxella devoid of the Tbp2 protein of the Moraxella strain and any other protein of the Moraxella strain and recombinantly-produced Tbp2 protein of a strain of Moraxella devoid of the Tbpl protein of the Moraxella strain and any other protein of the Moraxella strain and any other protein of the Moraxella strain. The Moraxella strain may be M. catarrhalis 4223 strain, M. catarrhalis Q8 strain or M. catarrhalis R1 strain.

In accordance with another aspect of the invention, an immunogenic composition is provided which comprises at least one active component selected from at least one nucleic acid molecule as provided herein and at least one recombinant protein as provided herein, and a pharmaceutically acceptable carrier therefor or vector therefor. The at least one active component produces an immune response when administered to a host.

The immunogenic compositions provided herein may be formulated as vaccines for in vivo administration to a For such purpose, the compositions formulated as a microparticle, capsule, ISCOM liposome preparation. The immunogenic composition may be provided in combination with a targeting molecule for delivery to specific cells of the immune system or to mucosal surfaces. The immunogenic compositions of the invention (including vaccines) may further comprise at one other immunogenic or immunostimulating material and the immunostimulating material may be at least one adjuvant or at least one cytokine. Suitable adjuvants for use in the present invention include (but are not limited to) aluminum phosphate, aluminum

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hydroxide, QS21, Quil A, derivatives and components ISCOM matrix, calcium phosphate, thereof. zinc hydroxide, a glycolipid analog, hydroxide, octadecyl ester of an amino acid, a muramyl dipeptide polyphosphazene, ISCOPREP, DC-chol, DDBA Advantageous combinations of adjuvants are lipoprotein. described in copending United States Patent Applications 08/261,194 filed June 16, 1994 and 08/483,856, filed June 7, 1995, assigned to the assignee hereof and the disclosures of which are incorporated herein by reference thereto (WO 95/34308).

In accordance with another aspect of the invention, there is provided a method for generating an immune response in a host, comprising the step of administering to a susceptible host, such as a human, an effective amount of the immunogenic composition provided herein. The immune response may be a humoral or a cell-mediated immune response and may provide protection against disease caused by Moraxella. Hosts in which protection against disease may be conferred include primates, including humans.

In a further aspect, there is provided a live vector for delivery of transferrin receptor to a host, comprising a vector containing the nucleic acid molecule as described above. The vector may be selected from Salmonella, BCG, adenovirus, poxvirus, vaccinia and poliovirus.

The nucleic acid molecules provided herein are useful in diagnostic applications. Accordingly, in a further aspect of the invention, there is provided a method of determining the presence, in a sample, of nucleic acid encoding a transferrin receptor protein of a strain of *Moraxella*, comprising the steps of:

(a) contacting the sample with a nucleic acid molecule as provided herein to produce duplexes comprising the nucleic acid molecule and any nucleic

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acid molecule encoding the transferrin receptor protein of a strain of *Moraxella* present in the sample and specifically hybridizable therewith; and

- (b) determining the production of the duplexes.
- In addition, the present invention provides a diagnostic kit for determining the presence, in a sample, of nucleic acid encoding a transferrin receptor protein of a strain of *Moraxella*, comprising:
 - (a) a nucleic acid molecule as provided herein;
- (b) means for contacting the nucleic acid molecule with the sample to produce duplexes comprising the nucleic acid molecule and any such nucleic acid present in the sample and hybridizable with the nucleic acid molecule; and
- (c) means for determining production of the duplexes.

The invention further includes the use of the nucleic acid molecules and proteins provided herein as medicines. The invention additionally includes the use of the nucleic acid molecules and proteins provided herein in the manufacture of medicaments for protection against infection by strains of *Moraxella*.

Advantages of the present invention include:

- an isolated and purified nucleic acid molecule encoding a transferrin receptor protein of a strain of *Moraxella* or a fragment or an analog of the transferrin receptor protein;
- recombinantly-produced transferrin receptor proteins, including Tbpl and Tbp2, free from each other and other *Moraxella* proteins; and
- diagnostic kits and immunological reagents for specific identification of *Moraxella*.

BRIEF DESCRIPTION OF DRAWINGS

The present invention will be further understood from the following description with reference to the

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drawings, in which:

Figure 1 shows the amino acid sequences (SEQ ID Nos: 17 and 18) of a conserved portion of Tbpl proteins used for synthesis of degenerate primers used for PCR amplification of a portion of the *M. catarrhalis* 4223 tbpA gene;

Figure 2 shows a restriction map of clone LEM3-24 containing the tbpA and tbpB genes from M. catarrhalis isolate 4223;

Figure 3 shows a restriction map of the tbpA gene for M. catarrhalis 4223;

Figure 4 shows a restriction map of the *tbpB* gene for *M. catarrhalis* 4223;

Figure 5 shows the nucleotide sequence of the *tbpA* gene (SEQ ID No: 1 - entire sequence and SEQ ID No: 2 - coding sequence) and the deduced amino acid sequence of the Tbpl protein from *M. catarrhalis* 4223 (SEQ ID No: 9 - full length and SEQ ID No: 10 - mature protein). The leader sequence (SEQ ID No: 19) is shown by underlining;

Figure 6 shows the nucleotide sequence of the *tbpB* gene (SEQ ID No: 3 - entire sequence and SEQ ID No: 4 - coding sequence) and the deduced amino acid sequence of the Tbp2 protein from *M. catarrhalis* 4223 (SEQ ID Nos: 11 - full length and SEQ ID No: 12 - mature protein).

The leader sequence (SEQ ID No: 20) is shown by underlining;

Figure 7 shows a restriction map of clone SLRD-A containing the tbpA and tbpB genes from M. catarrhalis Q8;

Figure 8 shows a restriction map of the tbpA gene from M. catarrhalis Q8;

Figure 9 shows a restriction map of the *tbpB* gene from *M. catarrhalis* Q8;

Figure 10 shows the nucleotide sequence of the tbpA gene (SEQ. ID No: 5 - entire sequence and SEQ ID No: 6 - coding sequence) and the deduced amino acid sequence of

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the Tbp1 protein from *M. catarrhalis* Q8 (SEQ ID No: 13 - full length and SEQ ID No: 14 - mature protein);

Figure 11 shows the nucleotide sequence of the tbpB gene (SEQ. ID No: 7 - entire sequence and SEQ ID No: 8 - coding sequence) and the deduced amino acid sequence of the Tbp2 protein from M. catarrhalis Q8 (SEQ ID No: 15 - full length and SEQ ID No: 16 - mature protein);

Figure 12 shows a comparison of the amino acid sequences of Tbp1 from *M. catarrhalis* strain 4223 (SEQ ID No: 9) and Q8 (SEQ ID No: 13), *H. influenzae* strain Eagan (SEQ ID No: 21), *N. meningitidis* strains B16B6 (SEQ ID No: 22) and M982 (SEQ ID No: 23), and *N. gonorrhoeae* strain FA19 (SEQ ID No: 24). Dots indicate identical residues and dashes have been inserted for maximum alignment;

Figure 13 shows a comparison of the amino acid sequences of Tbp2 from *M. catarrhalis* isolate 4223 (SEQ ID No: 11) and Q8 (SEQ ID No: 15), *H. influenzae* strain Eagan (SEQ ID No: 25), *N. meningitidis* strains B16B6 (SEQ ID No: 26) and M918 (SEQ ID No: 27), and *N. gonorrhoeae* strain FA19 (SEQ ID No: 28). Dots indicate identical residues and dashes have been inserted for maximum alignment;

Figure 14 shows the construction of plasmid pLEM29 for expression of recombinant Tbp1 protein from E. coli;

Figure 15 shows an SDS-PAGE analysis of the expression of Tbpl protein by *E. coli* cells transformed with plasmid pLEM29;

Figure 16 shows a flow chart for purification of recombinant Tbpl protein;

Figure 17 shows an SDS-PAGE analysis of purified recombinant Tbpl protein;

Figure 18 shows the construction of plasmid pLEM33 and pLEM37 for expression of TbpA gene from M. catarrhalis 4223 in E. coli without and with a leader sequence respectively;

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Figure 19 shows an SDS-PAGE analysis of the expression of rTbp2 protein by $E.\ coli$ cells transformed with plasmid pLEM37;

Figure 20 shows the construction of plasmid sLRD35B for expression of the *tbpB* gene from *M.* catarrhalis Q8 in *E. coli* without a leader sequence, and the construction of plasmid SLRD35A for expression of the *tbpB* gene from *M. catarrhalis* Q8 in *E. coli* with a leader sequence. Restriction site B = BamHI; Bg = Bgl II; H = Hind III; R = EcoRI;

Figure 21 shows SDS PAGE analysis of the expression of rTbp2 protein in *E. coli* cells, transformed with plasmids SLRD35A and SLRD35B;

Figure 22 shows a flow chart for purification of recombinant Tbp2 protein from *E. coli*;

Figure 23, which includes Panels A and B, shows an SDS-PAGE analysis of the purification of recombinant Tbp2 protein from *M. catarrhalis* strains 4223 (Panel A) and Q8 (Panel B) from expression in *E. coli*;

Figure 24 shows the binding of Tbp2 to human transferrin;

Figure 25, which includes Panels A, B and C, shows the antigenic conservation of Tbp2 protein amongst strains of *M. catarrhalis*;

Figure 26 shows a restriction map of the tbpB gene for M. catarrhalis R1;

Figure 27 shows the nucleotide sequence of the *tbpB* gene (SEQ ID No: 45 - entire sequence and SEQ ID No: 46 - coding sequence) and the deduced amino acid sequence of the Tbp2 protein of *M. catarrhalis* R1 (SEQ ID No: 47); and

Figure 28 shows a comparison of the amino acid sequences of Tbp2 for *M. catarrhalis* 4223 (SEQ ID No: 21), Q8 (SEQ ID No: 15) and R1 (SEQ ID No: 47). Dots indicate identical residues and dashes have been inserted for maximum alignment. The asterisks indicate

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stop codons.

GENERAL DESCRIPTION OF THE INVENTION

Any Moraxella strain may be conveniently used to provide the purified and isolated nucleic acid, which may be in the form of DNA molecules, comprising at least a portion of the nucleic acid coding for a transferrin receptor as typified by embodiments of the present invention. Such strains are generally available from clinical sources and from bacterial culture collections, such as the American Type Culture Collection.

this application, the terms "transferrin receptor" (TfR) and "transferrin binding proteins" (Tbp) are used to define a family of Tbpl and/or Tbp2 proteins which includes those having variations in their amino acid sequences including those naturally occurring in strains of, for example, Moraxella. purified and isolated DNA molecules comprising at least a portion coding for transferrin receptor of the present invention also include those encoding functional analogs of transferrin receptor proteins Tbp1 and Moraxella. In this application, a first protein is a "functional analog" of a second protein if the first protein is immunologically related to and/or has the same function as the second protein. The functional analog may be, for example, a fragment of the protein, or a substitution, addition or deletion mutant thereof.

Chromosomal from M. DNA catarrhalis digested with Sau3A in order to generate fragments within a 15 to 23 kb size range, and cloned into the BamHI site of the lambda vector EMBL3. The library was screened with anti-Tbpl guinea pig antisera, positive clone LEM3-24, containing an insert approximately 13.2 kb in size was selected for further analysis. Lysate from E. coli LE392 infected with LEM3-24 was found to contain a protein approximately 115 kDa

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in size, which reacted on Western blots with anti-Tbpl antisera. A second protein, approximately 80 kDa in size, reacted with the anti-Tbp2 guinea pig antisera on Western blots.

In order to localize the tbpA gene on the 13.2 kb insert of LEM3-24, degenerate PCR primers were used to amplify a small region of the putative tbpA gene of M. catarrhalis 4223. The sequences of the degenerate oligonucleotide primers were based upon conserved amino acid sequences within the Tbpl proteins of Neisseria and Haemophilus species and are shown Figure 1 (SEQ ID Nos: 17 and 18). A 300 base-pair amplified product was generated and its location within the 4223 tbpA gene is indicated by bold letters in Figure 5 (SEQ ID No: 29). The amplified product was subcloned into the vector pCRII, labelled, and used to а Southern blot containing restrictionendonuclease digested clone LEM3-24 DNA. The probe hybridized to a 3.8 kb HindIII-HindIII, a 2.0 kb AvrII-AvrII, and 4.2 kb SalI-SphI fragments (Figure 2).

The 3.8 kb HindIII-HindIII fragment was subcloned into pACYC177, and sequenced. A large open reading frame was identified, and subsequently found to contain approximately 2 kb of the putative tbpA gene. remaining 1 kb of the tbpA gene was obtained subcloning an adjacent downstream HindIII-HindIII fragment into vector pACYC177. The nucleotide sequence of the tbpA gene from M. catarrhalis 4223 (SEQ ID Nos: 1 and 2), and the deduced amino acid sequence (SEQ ID No: 9 - full length; SEQ ID No: 10 mature protein) are shown in Figure 5.

Chromosomal DNA from *M. catarrhalis* strain Q8 was digested with Sau3A I and 15-23 kb fragments were ligated with BamHI arms of EMBL3. A high titre library was generated in *E. coli* LE392 cells and was screened using oligonucleotide probes based on the 4223 *tbpA*

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sequence. Phage DNA was prepared and restriction enzyme analysis revealed that inserts of about 13-15 kb had been cloned. Phage clone SLRD-A was used to subclone fragments for sequence analysis. A cloning vector (pSKMA) was generated to facilitate cloning of the fragments and plasmids pSLRD1, pSLRD2, pSLRD3, pSLRD4 and pSLRD5 were generated which contain all of tbpA and most of tbpB. The nucleotide (SEQ ID Nos: 5 and 6) and deduced amino acid sequence (SEQ ID No: 13 - full length, SEQ ID No: 14 - mature protein) of the tbpA gene from strain Q8 are shown in Figure 10.

The deduced amino acid sequences for the Tbpl protein encoded by the *tbpA* genes were found to share some nomology with the amino acid sequences encoded by genes from a number of *Neisseria* and *Haemophilus* species (Figure 12; SEQ ID Nos: 21, 22, 23 and 24).

Prior to the present discovery, tbpA identified in species of Neisseria, Haemophilus, Actinobacillus have been found to be preceded by a tbpB gene with several conserved regions. The two genes typically are separated by a short intergenic sequence. However, a tbpB gene was not found upstream of the tbpA gene in M. catarrhalis 4223. In order to localize the tbpB gene within the 13.2 kb insert of clone LEM3-24, a denerate oligonucleotide probe was synthesized based upon an amino acid sequence EGGFYGP (SEQ ID No: 30), conserved among Tbp2 proteins of several species. oligonucleotide was labelled and used to Southern blot containing different restriction endonuclease fragments of clone LEM3-24. The probe hybridized to a 5.5 kb NheI-SalI fragment, subsequently was subcloned into pBR328, and sequenced. The fragment contained most of the putative tbpB gene, with the exception of the promoter region. The clone LEM3-24 was sequenced to obtain the remaining upstream sequence. The tbpB gene was located approximately 3 kb

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downstream from the end of the tbpA gene, in contrast to the genetic organization of the tbpA and tbpB genes in Haemophilus and Neisseria. The nucleotide sequence (SEQ ID Nos: 3 and 4) of the tbpB gene from M. catarrhalis 4223 and the deduced amino acid sequence (SEQ ID Nos: 11, 12) are shown in Figure 6. The tbpB gene from M. catarrhalis Q8 was also cloned and sequenced. nucleotide sequence (SEQ ID Nos: 7 and 8) deduced amino acid sequence (SEQ ID Nos: 15 and 16) are shown in Figure 11. The tbpB gene from M. catarrhalis was also cloned and sequenced. The nucleotide sequence (SEQ ID Nos: 45 and 46) and the deduced amino acid sequence (SEQ ID No: 47) are shown in Figure 27. Regions of homology are evident between catarrhalis Tbp2 amino acid sequences as shown in the comparative alignment of Figure 28 (SEQ ID Nos: 11, 15 and 47) and between the M. catarrhalis Tbp2 amino acid sequences and the Tbp2 sequences of a number Neisseria and Haemophilus species, as shown in the comparative alignment in Figure 13 (SEQ ID Nos: 25, 26, 27, 28).

Cloned tbpA and tbpB genes were expressed in E. coli to produce recombinant Tbpl and Tbp2 proteins free of other Moraxella proteins. These recombinant proteins were purified and used for immunization.

The antigenic conservation of Tbp2 protein amongst strains of *M. catarrhalis* was demonstrated by separation of the proteins in whole cell lysates of *M. catarrhalis* or strains of *E. coli* expressing recombinant Tbp2 proteins by SDS PAGE and antiserum immunoblotting with anti-4223 rTbp2 antiserum or anti-Q8 rTbp2 antiserum raised in guinea pigs. *M. catarrhalis* strains 3, 56, 135, 585, 4223, 5191, 8185 and ATCC 25240 were tested in this way and all showed specific reactivity with anti-4223 rTbp2 or anti-Q8 rTbp2 antibody (Figure 25).

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In addition, the ability of anti-rTbp2 antibodies from one strain to recognize native or recombinant protein from the homologous or heterologous strain by ELISA is shown in Table 1 below.

Amino acid sequencing of the N-termini and cyanogen bromide fragments of transferrin receptor from *M. catarrhalis* 4223 was undertaken. Both N-termini of Tbpl and Tbp2 were blocked. The putative signal sequences of Tbpl and Tbp2 are indicated by underlining in Figures 5 and 6 (SEQ ID Nos: 19 and 20) respectively. The deduced amino acid sequences for the N-terminal region of Tbp2 suggests a lipoprotein structure.

Results shown in Tables 1 and 2 below illustrate the ability of anti-Tbpl and anti-Tbp2 guinea pig antisera, produced by the immunization with Tbpl or Tbp2, to lyze M. catarrhalis. The results show that the antisera produced by immunization with Tbpl or Tbp2 protein isolated from M. catarrhalis isolate 4223 were bactericidal against a homologous non-clumping catarrhalis strain RH408 (a strain previously deposited in connection with United States Patent Application No. 08/328,589, assigned to the assignee hereof, 96/12733) with the American Type Culture Collection, located at 1301 Parklawn Drive, Rockville, Maryland 20852, USA under the terms of the Budapest Treaty on December 13, 1994 under ATCC Deposit No. 55,637) derived from isolate 4223. In addition, antisera produced by immunization with Tbp1 protein isolated from Μ. catarrhalis 4223 bactericidal were against the heterologous non-clumping strain Q8 (a gift from Dr. M.G. Bergeron, Centre Hospitalier de l'Université Laval, St. Foy, Quebec). In addition, antiserum raised against recombinant Tbp2 (rTbp2) protein was bacteriacidal against the homologous strain of M. catarrhalis.

The ability of isolated and purified transferrin binding proteins to generate bactericidal antibodies is

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in vivo evidence of utility of these proteins as vaccines to protect against disease caused by Moraxella.

Thus, in accordance with another aspect of the present invention, there is provided a vaccine against infection caused by *Moraxella* strains, comprising an immunogenically-effective amount of a transferrin binding protein from a strain of *Moraxella* and a physiologically-acceptable carrier therefor. Vaccine preparations may comprise antigenically or sequence divergent transferrin binding proteins.

The transferrin binding protein provided herein is useful as a diagnostic reagent, as an antigen for the generation of anti-transferrin protein binding antibodies, as an antigen for vaccination against the disease caused by species of *Moraxella* and for detecting infection by *Moraxella* and other such bacteria.

The transferrin binding protein provided herein may also be used as a carrier protein for haptens, polysaccharides or peptides to make conjugate vaccines against antigenic determinants unrelated to transferrin binding proteins. In additional embodiments of the present invention, therefore, the transferrin binding protein as provided herein may be used as a carrier molecule to prepare chimeric molecules and conjugate vaccines (including glycoconjugates) against pathogenic bacteria, including encapsulated bacteria. Thus, for example, glycoconjugates of the present invention may be used to confer protection against disease and infection caused by any bacteria having polysaccharide antigens including lipooligosaccharides (LOS) and PRP. bacterial pathogens may include, for example, Haemophilus influenzae, Streptococcus pneumoniae, Escherichia coli, Neisseria meningitidis, Salmonella typhi, Streptococcus mutans, Cryptococcus neoformans, Klebsiella, Staphylococcus aureus and Pseudomonas aeruginosa. Particular antigens which can be conjugated

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to transferrin binding protein and methods to achieve such conjugations are described in U.S. Patent Application No. 08/433,522 filed November 23, 1993 (WO 94/12641), assigned to the assignee hereof and the disclosure of which is hereby incorporated by reference thereto.

In another embodiment, the carrier function of transferrin binding protein may be used, for example, to induce an immune response against abnormal polysaccharides of tumour cells, or to produce antitumour antibodies that can be conjugated to chemotherapeutic or bioactive agents.

The invention extends to transferrin binding proteins from Moraxella catarrhalis for use as an active ingredient in a vaccine against disease caused by infection with Moraxella. The invention also extends to a pharmaceutical vaccinal composition containing transferrin binding proteins from Moraxella catarrhalis and optionally, a pharmaceutically acceptable carrier and/or diluent.

In a further aspect the invention provides the use of transferrin binding proteins for the preparation of a pharmaceutical vaccinal composition for immunization against disease caused by infection with *Moraxella*.

It is clearly apparent to one skilled in the art, that the various embodiments of the present invention have many applications in the fields of vaccination, diagnosis, treatment of, for example, *Moraxella* infections and the generation of immunological and other diagnostic reagents. A further non-limiting discussion of such uses is further presented below.

1. Vaccine Preparation and Use

Immunogenic compositions, suitable to be used as vaccines, may be prepared from immunogenic transferrin receptor proteins, analogs and fragments thereof encoded by the nucleic acid molecules as well as the nucleic

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acid molecules disclosed herein. The vaccine elicits an immune response which produces antibodies, including anti-transferrin receptor antibodies and antibodies that are opsonizing or bactericidal. Should the vaccinated subject be challenged by *Moraxella*, the antibodies bind to the transferrin receptor and thereby prevent access of the bacteria to an iron source which is required for viability. Furthermore, opsonizing or bactericidal anti-transferrin receptor antibodies may also provide protection by alternative mechanisms.

Immunogenic compositions, including vaccines, may prepared as injectables, as liquid solutions emulsions. The transferrin receptor proteins, analogs fragments thereof and encoding nucleic molecules may be mixed with pharmaceutically acceptable excipients which are compatible with the transferrin receptor proteins, fragments, analogs or nucleic acid Such excipients may include water, saline, molecules. dextrose, glycerol, ethanol, and combinations thereof. The immunogenic compositions and vaccines may further contain auxiliary substances, such as emulsifying agents, pH buffering agents, or adjuvants, enhance the effectiveness the of vaccines. Immunogenic compositions and vaccines may be administered parenterally, by injection subcutaneously, intradermally or intramuscularly. Alternatively, the immunogenic compositions provided according to present invention, may be formulated and delivered in a manner to evoke an immune response at mucosal surfaces. Thus, the immunogenic composition may be administered to mucosal surfaces by, for example, the nasal or oral (intragastric) routes. The immunogenic composition may be provided in combination with a targeting molecule for delivery to specific cells of the immune system or to mucosal surfaces. Some such targeting molecules include vitamin B12 and fragments of bacterial toxins,

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described in WO 92/17167 (Biotech Australia Pty. Ltd.), and monoclonal antibodies, as described in U.S. Patent No. 5,194,254 (Barber et al). Alternatively, other modes of administration, including suppositories and oral formulations, may be desirable. For suppositories, binders and carriers for may include, example, polyalkalene glycols or triglycerides. formulations may include normally employed incipients pharmaceutical as, for example, grades saccharine, cellulose and magnesium carbonate. compositions may take the form solutions, of suspensions, tablets, pills, capsules, sustained release formulations or powders and contain about 1 to 95% of the transferrin receptor proteins, fragments, analogs and/or nucleic acid molecules.

vaccines are administered in compatible with the dosage formulation, and in such amount as will be therapeutically effective, protective and immunogenic. The quantity to be administered depends on the subject to be treated, including, for example, the capacity of the individual's immune system to synthesize antibodies, and, if needed, to produce a cell-mediated immune response. Precise amounts active ingredient required to be administered depend on the judgment of the practitioner. However, suitable dosage ranges are readily determinable by one skilled in the art and may be of the order of micrograms of the transferrin receptor proteins, analogs and fragments thereof and/or nucleic acid molecules. Suitable regimes for initial administration and booster doses are also variable, but may include an initial administration followed by subsequent administrations. The dosage of vaccine may also depend on the route administration and will vary according to the size of the host.

The nucleic acid molecules encoding the transferrin

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receptor of Moraxella may be used directly immunization by administration of the DNA directly, for example, by injection for genetic immunization or by constructing a live vector, such as Salmonella, BCG, adenovirus, poxvirus, vaccinia or poliovirus containing the nucleic acid molecules. A discussion of some live vectors that have been used to carry heterologous antigens to the immune system is contained in, example, O'Hagan (ref 22). Processes for the direct injection of DNA into test subjects for immunization are described in, for example, Ulmer et al. (ref. 23).

Immunogenicity can be significantly improved if the antigens are co-administered with adjuvants, commonly used as an 0.05 to 1.0 percent solution in phosphate buffered saline. Adjuvants enhance the immunogenicity an antigen but are not necessarily immunogenic themselves. Adjuvants may act by retaining the antigen locally near the site of administration to produce a depot effect facilitating a slow, sustained release of antigen to cells of the immune system. Adjuvants can also attract cells of the immune system to an antigen depot and stimulate such cells to elicit immune responses.

Immunostimulatory agents or adjuvants have been 25 used for many years to improve the host immune responses to, for example, vaccines. Intrinsic adjuvants, such as lipopolysaccharides, normally are the components of killed or attenuated bacteria used as vaccines. 30 Extrinsic adjuvants are immunomodulators typically non-covalently linked to antigens and are formulated to enhance the host immune responses. adjuvants have been identified that enhance the immune response to antigens delivered parenterally. 35 these adjuvants are toxic, however, and can undesirable side-effects, making them unsuitable for use

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in humans and many animals. Indeed, only aluminum hydroxide and aluminum phosphate (collectively commonly referred to as alum) are routinely used as adjuvants in human and veterinary vaccines. The efficacy of alum in increasing antibody responses to diphtheria and tetanus toxoids is well established and an HBsAg vaccine has been adjuvanted with alum. While the usefulness of alum is well established for some applications, limitations. For example, alum is ineffective for influenza vaccination and inconsistently elicits a cell mediated immune response. The antibodies elicited by alum-adjuvanted antigens are mainly of the IgGl isotype in the mouse, which may not be optimal for protection by some vaccinal agents.

A wide range of extrinsic adjuvants can provoke potent immune responses to antigens. These include saponins complexed to membrane protein antigens (immune stimulating complexes), pluronic polymers with mineral oil, killed mycobacteria and mineral oil, Freund's complete adjuvant, bacterial products, such as muramyl dipeptide (MDP) and lipopolysaccharide (LPS), as well as lipid A, and liposomes.

To efficiently induce humoral immune responses (HIR) and cell-mediated immunity (CMI), immunogens are often emulsified in adjuvants. Many adjuvants are inducing granulomas, acute and chronic inflammations (Freund's complete adjuvant, FCA), cytolysis (saponins and pluronic polymers) pyrogenicity, arthritis and anterior uveitis (LPS and MDP). Although FCA is an excellent adjuvant and widely used in research, it is not licensed for use in human or veterinary vaccines because of its toxicity.

Desirable characteristics of ideal adjuvants include:

- 35 (1) lack of toxicity;
 - (2) ability to stimulate a long-lasting immune

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response;

- (3) simplicity of manufacture and stability in long-term storage;
- (4) ability to elicit both CMI and HIR to antigens administered by various routes, if required;
- (5) synergy with other adjuvants;
- (6) capability of selectively interacting with populations of antigen presenting cells (APC);
- (7) ability to specifically elicit appropriate $T_{H}1$ or $T_{H}2$ cell-specific immune responses; and
 - (8) ability to selectively increase appropriate antibody isotype levels (for example, IgA) against antigens.
- U.S. Patent No. 4,855,283 granted to Lockhoff et al on August 8, 1989, which is incorporated herein by 15 reference thereto, teaches glycolipid analogues including N-glycosylamides, N-glycosylureas glycosylcarbamates, each of which is substituted in the sugar residue by an amino acid, as immuno-modulators or 20 adjuvants. Thus, Lockhoff et al. 1991 (ref. reported that N-glycolipid analogs displaying structural similarities to the naturally-occurring glycolipids, such as glycophospholipids and glycoglycerolipids, are capable of eliciting strong immune responses in both 25 herpes simplex virus vaccine and pseudorabies virus Some glycolipids have been synthesized from long chain-alkylamines and fatty acids that are linked directly with the sugars through the anomeric carbon atom, to mimic the functions of the naturally occurring 30 lipid residues.
 - U.S. Patent No. 4,258,029 granted to Moloney, assigned to the assignee hereof and incorporated herein by reference thereto, teaches that octadecyl tyrosine hydrochloride (OTH) functions as an adjuvant when complexed with tetanus toxoid and formalin inactivated type I, II and III poliomyelitis virus vaccine. Also,

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Nixon-George et al. 1990, (ref. 25) reported that octadecyl esters of aromatic amino acids complexed with a recombinant hepatitis B surface antigen, enhanced the host immune responses against hepatitis B virus.

2. Immunoassays

The transferrin receptor proteins, analogs and/or fragments thereof of the present invention are useful as immunogens, as antigens in immunoassays including enzyme-linked immunosorbent assays (ELISA), RIAs and other non-enzyme linked antibody binding assays procedures known in the art for the detection of anti-Moraxella, transferrin receptor protein antibodies. ELISA assays, the transferrin receptor protein, analogs and/or fragments corresponding to portions of protein, are immobilized onto a selected surface, example, a surface capable of binding proteins peptides such as the wells of a polystyrene microtiter After washing to remove incompletely adsorbed transferrin receptor, analogs and/or fragments, a nonspecific protein such as a solution of bovine serum (BSA) or casein that is known antigenically neutral with regard to the test sample may be bound to the selected surface. This allows blocking of nonspecific adsorption sites the immobilizing surface and thus reduces the background caused by non-specific bindings of antisera onto the surface.

The immobilizing surface is then contacted with a sample, such as clinical or biological materials, to be tested in a manner conducive to immune complex (antigen/antibody) formation. This procedure include diluting the sample with diluents, such as BSA, bovine gamma globulin (BGG) and/or phosphate buffered saline (PBS)/Tween. The sample is then allowed to incubate for from about 2 to 4 hours, at temperatures such as of the order of about 25° to 37°C. Following

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incubation, the sample-contacted surface is washed to remove non-immunocomplexed material. The washing procedure may include washing with a solution such as PBS/Tween or a borate buffer.

Following formation of specific immunocomplexes the test sample and the bound transferrin between protein, analogs and/or fragments subsequent washing, the occurrence, and even amount, of immunocomplex formation may be determined by subjecting the immunocomplex to а second antibody specificity for the first antibody. If the test sample is of human origin, the second antibody is an antibody having specificity for human immunoglobulins and general IgG. To provide detecting means, the second antibody may have an associated activity such as an enzymatic activity that will generate, for example, a color development upon incubating with an appropriate chromogenic substrate. Quantification may then achieved by measuring the degree of color generation using, for example, a spectrophotometer.

3. Use of Sequences as Hybridization Probes

The nucleotide sequences of the present invention, comprising the sequence of the transferrin receptor gene, now allow for the identification and cloning of the transferrin receptor genes from any species of Moraxella.

The nucleotide sequences comprising the sequence of the transferrin receptor genes of the present invention are useful for their ability to selectively form duplex molecules with complementary stretches of other TfR genes. Depending on the application, a variety of hybridization conditions may be employed to achieve varying degrees of selectivity of the probe toward the other TfR genes. For a high degree of selectivity, relatively stringent conditions are used to form the duplexes, such as low salt and/or high temperature

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conditions, such as provided by 0.02 M to 0.15 M NaCl at temperatures of between about 50°C to 70°C. applications, less stringent hybridization conditions are required such as 0.15 M to 0.9 M salt, temperatures ranging from between about 20°C to 55°C. Hybridization conditions can also be rendered more stringent by the addition of increasing amounts of formamide, to destabilize the hybrid duplex. particular hybridization conditions can bе readily manipulated, and will generally be a method of choice depending on the desired results. In general, convenient hybridization temperatures in the presence of 50% formamide are: 42°C for a probe which is 95 to 100% homologous to the target fragment, 37°C for 90 to 95% homology and 32°C for 85 to 90% homology.

In a clinical diagnostic embodiment, the nucleic acid sequences of the TfR genes of the present invention may be used in combination with an appropriate means, such as a label, for determining hybridization. variety of appropriate indicator means are known in the art, including radioactive, enzymatic or other ligands, such as avidin/biotin and digoxigenin-labelling, which are capable of providing a detectable signal. In some diagnostic embodiments, an enzyme tag such as urease, alkaline phosphatase or peroxidase, instead radioactive tag may be used. In the case of enzyme tags, colorimetric indicator substrates are known which can be employed to provide a means visible to the human eye or spectrophotometrically, to identify specific hybridization with samples containing TfR sequences.

The nucleic acid sequences of TfR genes of the present invention are useful as hybridization probes in solution hybridizations and in embodiments employing solid-phase procedures. In embodiments involving solid-

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phase procedures, the test DNA (or RNA) from samples, such as clinical samples, including exudates, fluids (e. g., serum, amniotic fluid, middle effusion, sputum, bronchoalveolar lavage fluid) or even tissues, is adsorbed or otherwise affixed to a selected matrix or surface. The fixed, single-stranded nucleic acid is then subjected to specific hybridization with selected probes comprising the nucleic acid sequences of TfR genes or fragments thereof of the present invention under desired conditions. The selected conditions will depend on the particular circumstances based on the particular criteria required depending on, for example, the G+C contents, type of target nucleic acid, source of nucleic acid, size of hybridization probe etc. Following washing of the hybridization surface so as to remove non-specifically bound probe molecules, specific hybridization is detected, or even quantified, by means of the label. It is preferred to select nucleic acid sequence portions which conserved among species of Moraxella. The selected probe may be at least 18bp and may be in the range of about 30 to 90 bp.

4. Expression of the Transferrin Receptor Genes

Plasmid vectors containing replicon and control sequences which are derived from species compatible with the host cell may be used for the expression of the transferrin receptor genes in expression systems. The vector ordinarily carries a replication site, as well as marking sequences which are capable of providing phenotypic selection in transformed cells. For example, E. coli may be transformed using pBR322 which contains genes for ampicillin and tetracycline resistance and thus provides easy means for identifying transformed cells. The pBR322 plasmid, or other microbial plasmid or phage, must also contain, or be modified to contain, promoters which can be used by the host cell for

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expression of its own proteins.

In addition, phage vectors containing replicon and control sequences that are compatible with the host can be used as a transforming vector in connection with these hosts. For example, the phage in lambda $GEM^{TM}-11$ may be utilized in making recombinant phage vectors which can be used to transform host cells, such as E. coli LE392.

Promoters commonly used in recombinant DNA construction include the β -lactamase (penicillinase) and lactose promoter systems and other microbial promoters. such as the T7 promoter system as described in U.S. Patent No. 4,952,496. Details concerning the nucleotide sequences of promoters are known, enabling a skilled worker to ligate them functionally with genes. particular promoter used will generally be a matter of choice depending upon the desired results. Hosts that appropriate for expression of the transferrin receptor genes, fragments, analogs or variants thereof, may include E. coli, Bacillus species, Haemophilus, fungi, yeast, Moraxella, Bordetella, or the baculovirus expression system may be used.

In accordance with this invention, it is preferred to make the transferrin receptor protein, fragment or analog thereof, by recombinant methods, particularly since the naturally occurring TfR protein as purified from a culture of a species of Moraxella may include trace amounts of toxic materials or other contaminants. This problem can be avoided by using recombinantly produced TfR protein in heterologous systems which can be isolated from the host in a manner to minimize contaminants in the purified material. Particularly desirable hosts for expression in this regard include Gram positive bacteria which do not have LPS and are, therefore, endotoxin free. Such hosts include species of Bacillus and may be particularly useful for the

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production of non-pyrogenic transferrin receptor, fragments or analogs thereof. Furthermore, recombinant methods of production permit the manufacture of Tbp1 or Tbp2 respective analogs oror fragments separate from one another which is distinct from the normal combined proteins present in Moraxella.

Biological Deposits

Certain vectors that contain at least a portion coding for a transferrin receptor protein from strains 10 of Moraxella catarrhalis strain 4223 and Q8 and a strain of M. catarrhalis RH408 that are described and referred to herein have been deposited with the American Type Culture Collection (ATCC) located at 12301 Drive, Rockville, Maryland, USA, pursuant to the Budapest Treaty and prior to the filing of application. Samples of the deposited vectors bacterial strain will become available to the public and the restrictions imposed on access to the deposits will be removed upon grant of a patent based upon this United States patent application. In addition, the deposit will be replaced if viable samples cannot be dispensed by the Depository. The invention described and claimed herein is not to be limited in scope by the biological materials deposited, since the deposited embodiment is intended only as an illustration of the invention. equivalent or similar vectors or strains that encode similar or equivalent antigens as described in this application are within the scope of the invention.

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DEPOSIT	ATCC DESIGNATION	DATE DEPOSITED
Phage LEM3-24	97,381	December 4, 1995
Phage SLRD-A	97,380	December 4, 1995
Plasmid pLEM29	97,461	March 8, 1996
Plasmid pSLRD35A	97,833	January 13, 1997
Plasmid pLEM37	97,834	January 13, 1997
Strain RH408	55,637	December 9, 1994

EXAMPLES

above disclosure generally describes The present invention. A more complete understanding can be obtained by reference to following the specific These Examples are described solely for purposes of illustration and are not intended to limit the scope of the invention. Changes in form and substitution of equivalents contemplated as are circumstances may suggest or render expedient. Although specific terms have been employed herein, such terms are intended in a descriptive sense and not for purposes of limitations.

Methods of molecular genetics, protein biochemistry and immunology used but not explicitly described in this disclosure and these Examples are amply reported in the scientific literature and are well within the ability of those skilled in the art.

Example 1

This Example illustrates the preparation and immunization of guinea pigs with Tbpl and Tbp2 proteins from M. catarrhalis.

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Tbpl and Tbp2 proteins were obtained as follows:

Iron-starved crude total membrane preparations were diluted to 4 mg protein/ml in 50 mM Tris.HCl-1M NaCl, pH a total volume of 384 ml. Membranes were solubilized by the addition of 8 ml each of 0.5M EDTA and 30% sarkosyl and samples were incubated for 2 hours at room temperature, with gentle agitation. Solubilized membranes were centrifuged at 10K rpm for 20 min. of apo-hTf-Sepharose 4B were added to the supernatant, and incubated for 2 hours at room temperature, gentle shaking. The mixture was added into a column. The column was washed with 50 ml of 50mM Tris.HCl-1 M NaCl-250mM quanidine hydrochloride, contaminating proteins. Tbp2 was eluted from the column by the addition of 100 ml of 1.5M quanidine hydrochloride. Tbpl was eluted by the addition of 100 ml of 3M guanidine hydrochloride. The first 20 fractions were dialyzed against 3 changes of 50 Tris.HCl, pH 8.0. Samples were stored at -20°C, dialyzed against ammonium bicarbonate and lyophilized.

Guinea River) pigs (Charles were intramuscularly on day +1 with a 10 μ g dose of Tbpl or Tbp2 emulsified in complete Freund's adjuvant. were boosted on days +14 and +29 with the same dose of protein emulsified in incomplete Freund's adjuvant. Blood samples were taken on day +42, and sera were used for analysis of bactericidal antibody activity. addition. all antisera were assessed by immunoblot analysis for reactivity with M. catarrhalis proteins.

The bactericidal antibody activity of guinea pig anti-M. catarrhalis 4223 Tbpl or Tbp2 antisera was determined as follows. A non-clumping M. catarrhalis strain RH408, derived from isolate 4223, was inoculated into 20 ml of BHI broth, and grown for 18 hr at 37°C, shaking at 170 rpm. One ml of this culture was used to

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inoculate 20 ml of BHI supplemented with 25 ethylenediamine-di-hydroxyphenylacetic acid The culture was grown to an OD,, of 0.5. cells were diluted 1:200,000 in 140 mM NaCl, 93mM NaHCO,, 2mM Na barbiturate, 4mM barbituric acid, 0.5mM MgCl,.6H,0, 0.4mM CaCl,.2H,0, pH 7.6 (Veronal buffer), containing 0.1% bovine serum albumin (VBS) and placed on Guinea pig anti-M. catarrhalis 4223 Tbpl or Tpb2 antisera, along with prebleed control antisera, were heated to 56°C for 30 min. to inactivate endogenous complement. Serial twofold dilutions of each antisera in VBS were added to the wells of a 96-well Nunclon microtitre plate (Nunc, Roskilde, Denmark). Dilutions started at 1:8, and were prepared to a final volume of 25 μL in each well. 25 μL of diluted bacterial cells added to each of the wells. A guinea complement (Biowhittaker, Walkersville, MD) was diluted 1:10 in VBS, and 25 μL portions were added to each well. The plates were incubated at 37°C for 60 min, gently shaking at 70 rpm on a rotary platform. $50 \mu L$ of each reaction mixture were plated onto Mueller (Becton-Dickinson, Cockeysville, MD) agar plates. plates were incubated at 37°C for 72 hr and the number of colonies per plate were counted. Bactericidal titres were assessed as the reciprocal of the highest dilution of antiserum capable of killing greater than 50% of bacteria compared with controls containing pre-immune Results shown in Table 1 below illustrate the ability of the anti-Tbpl and anti-Tbp2 guinea pig antisera to lyze M. catarrhalis.

Example 2

This Example illustrates the preparation of chromosomal DNA from M. catarrhalis strains 4223 and Q8.

M. catarrhalis isolate 4223 was inoculated into 100 ml of BHI broth, and incubated for 18 hr at 37°C with

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shaking. The cells were harvested by centrifugation at $10,000 \times g$ for 20 min. The pellet was used for extraction of M. catarrhalis 4223 chromosomal DNA.

The cell pellet was resuspended in 20 ml of 10 mM $\,$ Tris-HCl (pH 7.5)-1.0 mM EDTA (TE). Pronase and SDS were added to final concentrations of 500 μ g/ml and 1.0%, respectively, and the suspension was incubated at 37°C for 2 hr. After several sequential extractions with phenol. phenol:chloroform (1:1),chloroform:isoamyl alcohol (24:1), the aqueous extract was dialysed, at 4°C, against 1.0 M NaCl for 4 hr, and against TE (pH 7.5) for a further 48 hr with three buffer changes. Two volumes of ethanol were added to the dialysate, and the DNA was spooled onto a glass rod. The DNA was allowed to air-dry, and was dissolved in 3.0 ml of water. Concentration was estimated, by UV spectrophotometry, to be about 290 μ g/ml.

M. catarrhalis strain Q8 was grown in BHI broth as described in Example 1. Cells were pelleted from 50 ml of culture by centrifugation at 5000 rpm for 20 minutes, at 4°C . The cell pellet was resuspended in 10 ml of TE (10 mM Tris-HCl, 1 mM EDTA, pH 7.5) and proteinase K and SDS were added to final concentrations of 500 $\mu g/ml$ and 1%, respectively. The sample was incubated at 37° C for 4 hours until a clear lysate was obtained. The lysate was extracted twice with Tris-saturated phenol/chloroform (1:1), and twice with chloroform. The final aqueous phase was dialysed for 24 hours against 2 X 1000 ml of 1 M NaCl at 4°C , changing the buffer once, and for 24hours against 2 \times 1000 ml of TE at 4°C, changing the buffer once. The final dialysate was precipitated with two volume of 100% ethanol. The DNA was spooled, dried and resuspended in 5 to 10 ml of TE buffer.

Example 3

This Example illustrates the construction of M.

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catarrhalis chromosomal libraries in EMBL3.

series of Sau3A restriction digests chromosomal DNA, in final volumes of 10 µL each, were carried out in order to optimize the conditions necessary to generate maximal amounts of restriction fragments within a 15 to 23 kb size range. Using the optimized digestion conditions, a large-scale digestion was set up in a 100 μL volume, containing the following: 50 μ L of chromosomal DNA (290 μ g/ml), 33 μ L water, 10 μL 10X Sau3A buffer (New England Biolabs), 1.0 μL BSA (10 mg/ml, New England Biolabs), and 6.3 μ L Sau3A (0.04) U/µL). Following a 15 min. incubation at 37°C, the digestion was terminated by the addition of 10 μL of 100 mM Tris-HCl (pH 8.0)-10 mM EDTA-0.1% bromophenol blueglycerol (loading buffer). Digested electrophoresed through a 0.5% agarose gel in 40 mM Tris acetate-2 mM Na,EDTA.2H,0 (pH8.5) (TAE buffer) at 50 V for б hr. The region containing restriction fragments within a 15 to 23 kb molecular size range was excised from the gel, and placed into dialysis tubing containing 3.0 ml of TAE buffer. DNA was electroeluted from the gel fragment by applying a field strength of 1.0 V/cm for 18 hr. Electroeluted DNA was extracted once each phenol and phenol:chloroform (1:1).precipitated with ethanol. The dried DNA was dissolved in 5.0 µL water.

Size-fractionated chromosomal DNA was ligated with BamHI-digested EMBL3 arms (Promega), using T4 DNA ligase in a final volume of 9 μ L. The entire ligation mixture was packaged into lambda phage using a commercial packaging kit (Amersham), following manufacturer's instructions.

The packaged DNA library was amplified on solid media. 0.1 ml aliquots of *Escherichia coli* strain NM539 in 10 mM MgSO. $(OD_{240} = 0.5)$ were incubated at 37°C for 15

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min. with 15 to 25 μL of the packaged DNA library. Samples were mixed with 3 ml of 0.6% agarose containing 1.0% BBL trypticase peptone-0.5% NaCl (BBL top agarose), mixtures plated onto were 1.5% agar containing 1.0% BBL trypticase peptone-0.5% NaCl, incubated at 37°C for 18 hr. 3 ml quantities of 50 mM Tris-HCl (pH 7.5)-8 mM magnesium sulfate heptahydrate-100 mM NaCl-0.01% (w/v) gelatin (SM buffer) were added to each plate, and plates were left at 4°C for 7 hr. buffer containing phage was collected from the plates, pooled together, and stored in a screwcap tube at 4°C, with chloroform.

Chromosomal DNA from M. catarrhalis strain Q8 was digested with Sau3A I (0.1 unit/30 μg DNA) at 37°C for 30 minutes and size-fractionated on a 0.6% low melting point agarose gel. DNA fragments of 15-23 kb were excised and the DNA was electroeluted for 25 minutes in dialysis tubing containing TAE (40 mM Tris acetate pH 8.5, 2 mM EDTA) at 150 V. The DNA was extracted once phenol/chloroform (1:1), precipitated. resuspended in water. The DNA was ligated overnight with EMBL3 BamH I arms (Promega) and the ligation mixture was packaged using the Lambda in vitro packaging kit (Stratagene) and plated onto E. coli LE392 cells. The library was titrated and stored at 4°C in the presence of 0.3% chloroform.

Example 4

This Example illustrates screening of the M. catarrhalis libraries.

30 Ten μ L aliquots of phage stock from the EMBL3/4223 sample prepared in Example 3 above were combined each with 100 μ L of *E. coli* strain LE392 in 10 mM MgSO4 (OD_{2.0} = 0.5) (plating cells), and incubated at 37°C for 15 min. The samples were mixed with 3 ml each of BBL top agarose, and the mixtures were poured onto 1.5% agarose

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plates containing 1% bacto tryptone-0.5% bacto yeast extract-0.05% NaCl (LB agarose; Difco) and supplemented with 200 μM EDDA. The plates were incubated at 37°C for 18 hr. Plaques were lifted onto nitrocellulose filters (Amersham Hybond-C Extra) using a standard protocol, and the filters were immersed into 5% bovine serum albumin (BSA; Boehringer) in 20 mM Tris-HCl (pH 7.5)-150 mM NaCl (TBS) for 30 min at room temperature, or 4°C overnight. Filters were incubated for at least 1 hr at room temperature, or 18 hr at 4°C, in containing a 1/1000 dilution of guinea pig anti-M. catarrhalis 4223 Tbp1 antiserum. Following sequential 10 min. washes in TBS with 0.05% Tween 20 (TBS-Tween), filters were incubated for 30 min. at room temperature in TBS-Tween containing a 1/4000 dilution of recombinant Protein G labelled horseradish with peroxidase (rProtein G-HRP; Zymed). Filters were washed as above, and submerged into CN/DAB substrate solution (Pierce). Color development was arrested by immersing the filters into water. Positive plaques were cored from the plates, and each placed into 0.5 ml of SM buffer containing a few drops of chloroform. The screening procedure was repeated two more times, until 100% of the lifted plaques were positive using the guinea pig anti-M. catarrhalis 4223 Tbp1 antiserum.

The EMBL3/Q8 library was plated onto LE392 cells on YT plates using 0.7% top agar in YT as overlay. Plaques were lifted onto nitrocellulose filters and the filters were probed with oligonucleotide probes labelled with 32 P α -dCTP (Random Primed DNA labeling kit, Boehringer Mannheim). The pre-hybridization was performed in sodium chloride/sodium citrate (SSC) buffer (ref. 27) at 37° C for 1 hour and the hybridization was performed at 42° C overnight. The probes were based upon an internal sequence of 4223 tbpA:

IRDLTRYDPG

(Seq ID No. 31)

4236-RD 5' ATTCGAGACTTAACACGCTATGACCCTGGC 3'

(Seq ID No 32)

5 4237-RD 5' ATTCGTGATTTAACTCGCTATGACCCTGGT 3' (Seq ID No 33).

Putative plaques were re-plated and submitted to second and third rounds of screening using the same procedures. Phage clone SLRD-A was used to subclone the tfr genes for sequence analysis.

Example 5

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This Example illustrates immunoblot analysis of the phage lysates using anti-M. catarrhalis 4223 Tbp1 and Tbp2 antisera.

Proteins expressed by the phage eluants selected in Example 4 above were precipitated as follows. 60 μL of each phage eluant were combined with 200 μL E. coli LE392 plating cells, and incubated at 37°C for 15 min.

The mixture was inoculated into 10 ml of 1.0% NZamine A-20 0.5% NaCl-0.1% casamino acids-0.5% yeast extract-0.2% sulfate magnesium heptahydrate (NZCYM broth). supplemented with 200 mM EDDA, and grown at 37°C for 18hr, with shaking. DNAse was added to 1.0 ml of the culture, to a final concentration of 50 $\mu g/ml$, and the 25 sample was incubated at 37°C for 30

Trichloroacetic acid was added to a final concentration of 12.5%, and the mixture was left on ice for 15 min.

Proteins were pelleted by centrifugation at 13,000 x g for 10 min, and the pellet was washed with 1.0 ml of acetone. The pellet was air-dried and resuspended in 50 μ L 4% SDS-20 mM Tris- HCl (pH 8.0)-0.2 mM EDTA (lysis buffer).

Following SDS-PAGE electrophoresis through an 11.5% gel, the proteins were transferred to Immobilon-P

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filters (Millipore) at a constant voltage of 20 V for 18 Tris-HCl,220mM in 25mM glycine-20% (transfer buffer). Membranes were blocked in 5% BSA in TBS for 30 min. at room temperature. Blots were exposed either to guinea pig anti-M. catarrhalis 4223 Tbp1, or to guinea pig anti-M. catarrhalis 4223 Tbp2 antiserum, diluted 1/500 in TBS-Tween, for 2 hr temperature. Following three sequential 10 min. washes in TBS-Tween, membranes were incubated in TBS-Tween containing a 1/4000 dilution of rProtein G-HRP for 30 min. at room temperature. Membranes were washed as described above, and immersed into CN/DAB substrate Color development was arrested by immersing solution. blots into water.

Three EMBL3 phage clones expressed both a 115 kDa protein which reacted with anti-Tbpl antiserum, and an 80 kDa protein, which reacted with anti-Tbp2 antiserum on Western blots and were thus concluded to contain genes encoding the transferrin receptor proteins of Moraxella catarrhalis.

Example 6

This Example illustrates the subcloning of the M. catarrhalis 4223 Tbpl protein gene, tbpA.

Plate lysate cultures of the recombinant phage described in Example 5 were prepared by combining phage eluant and *E. coli* LE392 plating cells, to produce confluent lysis on LB agar plates. Phage DNA was extracted from the plate lysates using a Wizard Lambda Preps DNA Purification System (Promega), according to manufacturer's instructions.

The EMBL3 clone LM3-24 was found to contain a 13.2 kb insert, flanked by two SalI sites. A probe to a tbpA gene was prepared and consisted of a 300 base pair amplified product generated by PCR using two degenerate oligonucleotide primers corresponding to an amino acid sequence of part of the Tbpl protein (Figure 1). The

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primer sequences were based upon the amino sequences NEVTGLG (SEQ ID No: 17) and GAINEIE (SEQ ID No: 18), which had been found to be conserved among the deduced amino acid sequences from several different N. meningitidis and Haemophilus influenzae tbpA genes. amplified product was cloned into pCRII (Invitrogen, San and sequenced. The deduced amino acid sequence shared homology with other putative amino acid sequences derived from N. meningitidis and H. influenzae tbpA genes (Figure 12). The subclone was linearized with NotI (New England Biolabs), and labelled using a digoxigenin random-labelling kit (Boehringer Mannheim), according to manufacturer's instructions. concentration of the probe was estimated to be 2 $ng/\mu L$.

DNA from the phage clone was digested with HindIII, SalI/SphI, or SalI/AvrII, and electrophoresed AvrII. through a 0.8% agarose gel. DNA was transferred to a nylon membrane (Genescreen Plus, Dupont) using an LKB VacuGene XL vacuum transfer apparatus (Pharmacia). Following transfer, the blot was air-dried, and prehybridized in 5X SSC-0.1% N-lauroylsarcosine-0.02% sodium dodecyl sulfate-1.0% blocking reagent (Boehringer Mannheim) in 10 mM maleic acid-15 mM NaCl (pH 7.5) (prehybridization solution). Labelled probe was added to the pre-hybridization solution to a final concentration of 6 ng/ml, and the blot was incubated in the probe solution at 42°C for 18 hr. The blot was washed twice in 2X SSC-0.1% SDS, for 5 min. each at room temperature, then twice in 0.1% SSC-0.1% SDS for 15 min. each at 60°C. Following the washes, the membrane equilibrated in 100mM maleic acid-150 mM NaCl (pH 7.5) (buffer 1) for 1 min, then left in 1.0% blocking reagent (Boehringer Mannheim) in buffer 1 (buffer 2) for 60 min, at room temperature. The blot was exposed to anti-DIGalkaline phosphatase (Boehringer Mannheim) 1/5000 in buffer 2, for 30 min. at room temperature.

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Following two 15 min. washes in buffer 1, the blot was equilibrated in 100 mM Tris-HCl (pH 9.5), 100 mM NaCl, 50 mM MgCl, (buffer 3) for 2 min. The blot was wetted with Lumigen PPD substrate (Boehringer-Mannheim), diluted 1/100 in buffer 3, then wrapped in Saran wrap, and exposed to X-ray film for 30 min. The probe hybridized to a 3.8 kb HindIII-HindIII, a 2.0 kb AvrII-AvrII, and a 4.2 kb SalI-SphI fragment.

In order to subclone the 3.8 kb HindIII-HindIII fragment into pACYC177, phage DNA from the EMBL3 clone, and plasmid DNA from the vector pACYC177 (New England Biolabs), were digested with HindIII, and fractionated by electrophoresis on a 0.8% agarose gel. The 3.8 kb HindIII-HindIII phage DNA fragment, and the 3.9 HindIII-HindIII pACYC177 fragment, were excised from the gel and purified using a Geneclean kit (Bio 101 Inc., LaJolla, CA), according to manufacturer's directions. Purified insert and vector were ligated together using T4 DNA ligase (New England Biolabs), and transformed into E. coli HB101 (Gibco BRL). A Qiagen Plasmid Midi-Kit (Qiagen) was used to extract and purify sequencingquality DNA from one of the ampicillinresistant/kanamycin-sensitive transformants, which found to carry a 3.8 kb HindIII-HindIII insert. subclone was named pLEM3. As described in Example 7, subsequent sequencing revealed that pLEM3 contained the first about 2.0 kb of tbpA sequence (Figures 2 and 5).

In order to subclone the remaining 1 kb of the tbpA gene, a 1.6 kb HindIII-HindIII fragment was subcloned into pACYC177 as described above, and transformed by electroporation into E. coli HB101 (Gibco BRL). A Midi-Plasmid DNA kit (Qiagen) was used to extract plasmid DNA from a putative kanamycin-sensitive transformant carrying a plasmid with a 1.6 kb HindIII-HindIII insert. The subclone was termed pLEM25. As described in

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Example 7 below, sequencing revealed that pLEM25 contained the remaining 1 kb of the tbpA gene (Figure 2 and 5).

Example 7

This Example illustrates the subcloning of the M. catarrhalis 4223 tbpB gene.

described above. in all Neisseriae and Haemophilus species examined prior to the present invention, tbpB genes been found have immediately upstream of the tbpA genes which share homology with the tbpA gene of M. catarrhalis 4223. However, the sequence upstream of M. catarrhalis 4223 did not correspond with other sequences encoding tbpB.

In order to localize the tbpB gene within the EMBL3 phage clone, a Southern blot was carried out using a degenerate probe from a highly conserved amino acid within the Tbp2 protein. Α degenerate oligonucleotide probe, was designed corresponding to the sequence encoding EGGFYGP (SEQ ID No: 30), which is conserved within the Tbp2 protein in a variety of Neisseriae and Haemophilus species. The probe was labelled digoxigenin using an with oligonucleotide tailing (Boehringer Mannheim), kit following manufacturer's instructions. HindIII - digested EMBL3 clone DNA was fractionated through a 0.8% agarose gel, and transferred to a Geneclean Plus nylon membrane as described in Example 6. Following hybridization as described above, the membrane was washed twice in 2X SSC-0.1% SDS, for 5 min. each at room temperature, then twice in 0.1% SSC-0.1% SDS for 15 min. each, at 50°C. Extection of the labelled probe was carried out as The probe hybridized to a 5.5 kb Nheldescribed above. SalI fragment.

The 5.5 kb *NheI-Sal*I fragment was subcloned into pBR328 as follows. LEM3-24 DNA, and pBR328 DNA, were digested with *NheI-Sal*I, and electrophoresed through

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0.8% agarose. The 5.5 kb NheI-SalI fragment, and the 4.9 kb pBR328 NheI-SalI fragments were excised from the gel, and purified using a Geneclean kit as described in Example 6. The fragments were ligated together using T4 DNA ligase, and transformed into E. coli DH5. Plasmid DNA kit (Qiagen) was used to extract DNA from an ampicillin resistant / tetracycline sensitive clone containing a 5.5 kb NheI-SalI insert. This subclone was termed pLEM23. Sequencing revealed that pLEM23 contained 2 kb of the tbpB gene from M. catarrhalis 4223 (Figure 2).

Example 8:

This Example illustrates the subcloning of M. catarrhalis Q8 tfr genes.

The M. catarrhalis Q8 tfr genes were subcloned as Phage DNA was prepared from plates. the top agarose layer from three confluent plates was scraped into 9 ml of SM buffer (0.1 M NaCl, 0.2% MgSO4, 50 mM Tris-HCl, pH 7.6, 0.01% gelatin) and 100 μ l of chloroform was added. The mixture was vortexed for 10 sec, then incubated at room temperature for 2h. cell debris was removed by centrifugation at 8000 rpm for 15 min at 4°C in an SS34 rotor (Sorvall model RC5C). The phage was pelleted by centrifugation at 35,000 rpm in a 70.1 Ti rotor at 10° C for 2h (Beckman model L8-80) and was resuspended in 500 µl of SM buffer. The sample was incubated at 4°C overnight, then RNAse and DNAse were added to final concentrations of 40 $\mu g/ml$ and 10 μ g/ml, respectively and the mixture incubated at 37°C To the mixture were added 10 μl of 0.5 M EDTA and 5 μl of 10% SDS and the sample was incubated at 6°C for 15 min. The mixture was extracted twice with phenol/chloroform (1:1) and twice with chloroform and the DNA was precipitated by the addition of 2.5 volumes of absolute ethanol.

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partial restriction map was generated fragments were subcloned using the external Sal I sites from EMBL3 and internal AvrII or EcoR I sites as indicated in Figure 4. In order to facilitate the subcloning, plasmid pSKMA was constructed introduces a novel multiple cloning site into pBluescript.SK (Stratagene). Oligonucleotides were used to introduce restriction sites for Mst II, Sfi I, and Avr II between the Sal I and Hind III pBluescript.SK:

Sfi I

Sal I Cla I Mst II Avr II HindIII

15 4639-RD 5' TCGACGGTAT CGATGGCC TTAG GGGC CTAGGA 3' (SEQ ID No: 34)

4640-RD 3' GCCATA GCTACCGG AATC CCCG GATCCTTCGA (SEQ ID No: 35)

Plasmid pSLRD1 contains a ~1.5 kb Sal I-Avr II fragment cloned into pSKMA; plasmids pSLRD2 and pSLRD4 contain ~2 kb and 4 kb AvrII-AvrII fragments cloned into pSKMA, respectively and contain the complete tbpA gene. Plasmid pSLRD3 contains a ~2.3 kb AvrII-EcoR I fragment cloned into pSKMA and plasmid SLRD5 is a 22.7 kb EcoRI - EcoRI fragment cloned into pSKMA. These two clones contain the complete tbpB gene (Figure 7).

Example 9

This Example illustrates sequencing of the M. catarrhalis tbp genes.

Both strands of the *tbp* genes subcloned according to Examples 6 to 8 were sequenced using an Applied Biosystems DNA sequencer. The sequences of the *M. catarrhalis* 4223 and Q8 *tbpA* genes are shown in Figures 5 and 10 respectively. A derived amino acid sequence was compared with other Tbp1 amino acid sequences, including

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Neisseriae meningitidis, of Neisseriae gonorrhoeae, and Haemophilus influenzae (Figure 12). The sequence of the M. catarrhalis 4223 and Q8 tbpB genes are shown in Figures 6 and 11 respectively. In order to obtain sequence from the putative beginning of the tbpB gene of M. catarrhalis 4223, sequence data were obtained directly from the clone LEM3-24 DNA. sequence was verified by screening clone DS-1754-1. The of the translated tbpB genes catarrhalis 4223 and Q8 shared homology with deduced Tbp2 amino acid sequences of Neisseria meningitidis, Neisseria gonorrhoeae, and Haemophilus influenzae (Figure 13).

Example 10

This Example illustrates the generation of an expression vector to produce recombinant Tbpl protein. The construction scheme is shown in Figure 14.

Plasmid DNA from subclone pLEM3, prepared described in Example 6, was digested with HindIII and BglI to generate a 1.84 kb BglI-HindIII fragment, containing approximately two-thirds of the tbpA gene. was added to the digest to eliminate comigrating 1.89kb BglI-HindIII vector fragment. Ιn addition, plasmid DNA from the vector pT7-7 digested with NdeI and HindIII. To create the beginning of the tbpA gene, an oligonucleotide was synthesized based upon the first 61 bases of the tbpA gene to the Ball site; an NdeI site was incorporated into the 5' Purified insert, vector and oligonucleotide were end. ligated together using T4 ligase (New England Biolabs), and transformed into $E.\ coli$ DH5 $\alpha.$ DNA was purified of the 4.4 kb ampicillin-resistant transformants containing correct restriction (pLEM27).

Purified pLEM27 DNA was digested with HindIII, ligated to the 1.6 kb HindIII-HindIII insert fragment

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of pLEM25 prepared as described in Example 6, and transformed into $E.\ coli$ DH5 α . DNA was purified from an ampicillin-resistant transformant containing the correct restriction sites (pLEM29), and was transformed by electroporation into BL21 (DE3) (Novagen; Madison, WI) to produce $E.\ coli$ pLEM29B-1.

A single isolated transformed colony was used to inoculate 100 ml of YT broth containing ampicillin, and the culture was grown at overnight, shaking at 200 rpm. 200 μ l of the overnight culture were inoculated into 10 ml of YT broth containing $100\mu g/ml$ ampicillin, and the culture was grown at 37° C to an OD₅₇₈ of 0.35. The culture was induced by the addition of 30 μl of 100 mM IPTG, and the culture was grown at 37°C for an additional 3 One ml of culture was removed at the time of induction (t=0), and at t=1 hr and t=3 hrs. One ml samples were pelleted by centrifugation, resuspended in 4%SDS-20 mM Tris.Cl, pH 8-200 μM EDTA (lysis buffer). Samples were fractionated on an 11.5% SDS-PAGE gel, and transferred onto Immobilon filters (Amersham). Blots were developed using anti-Tbp1 (M. catarrhalis 4223) antiserum, diluted 1:1000, as primary antibody, and rproteinG conjugated horseradish peroxidase (Zymed) as the secondary antibody. A chemiluminescent substrate (Lumiglo; Kirkegaard and Perry Laboratories, Gaithersburg, was used for detection. Induced recombinant proteins were visible on the Coomassie-stained gels (Fig 15). anti-Tbpl (4223)antiserum recognized the recombinant proteins on Western blots.

Example 11

This Example illustrates the extraction and purification of recombinant Tbpl of *M. catarrhalis* 4223.

Recombinant Tbpl protein, which is contained in inclusion bodies, was purified from *E. coli* cells

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expressing the tbpA gene (Example 10), by a procedure as shown in Figure 16. E. coli cells from a 500 ml culture, prepared as described in Example 10, resuspended in 50 ml of 50 mM Tris-HCl, Hq containing 0.1 M NaCl and 5 mM AEBSF inhibitor), and disrupted by sonication (3 \times 10 min. 70% duty circle). The extract was centrifuged at $20,000 \times g$ for 30 min. and the resultant supernatant contained > 85% of the soluble proteins from E. coli was discarded.

The remaining pellet (Figure 16, PPT₁) was further extracted in 50 ml of 50 mM Tris, pH 8.0 containing 0.5% Triton X-100 and 10 mM EDTA. After centrifugation at $20,000 \times g$ for 30 min., the supernatant containing residual soluble proteins and the majority of the membrane proteins was discarded.

The remaining pellet (Figure 16, PPT_2) was further extracted in 50 ml of 50 mM Tris, pH 8.0 containing 2M urea and 5 mM dithiothroitol (DTT). After centrifugation at 20,000 x g for 30 min., the resultant pellet (Figure 16, PPT_3) obtained after the above extraction contained the purified inclusion bodies.

The Tbpl protein was solubilized from PPT3 in 50 mM Tris, pH 8.0, containing 6 M guanidine hydrochloride and DTT. After centrifugation, the resultant supernatant was further purified on a Superdex 200 gel filtration column equilibrated in 50 mM Tris, pH 8.0, containing 2M guanidine hydrochloride and 5 mM DTT. fractions were analyzed by SDS-PAGE and those containing purified Tbp1 were pooled. Triton X-100 was added to the pooled Tbpl fraction to a final concentration of 0.1%. The fraction was then dialyzed overnight at 4°C against 50 mM Tris, pH 8.0 and then centrifuged at 20,000 x g for 30 min. The protein remained soluble under these conditions and the purified Tbpl was stored The purification procedure shown in Figure at -20° C.

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16 produced Tbp1 protein that was at least 70% pure as determined by SDS-PAGE analysis (Figure 17).

Example 12

This Example illustrates the construction of an expression plasmid for rTbp2 of *M. catarrhalis* 4223 without a leader sequence.

The construction scheme for the plasmid expressing rTbp2 is shown in Figure 18. Oligonucleotides were used to construct the first approximately 58 bp of the M. catarrhalis 4223 tbpB gene encoding the mature protein. An NdeI site was incorporated into the 5' end of the oligonucleotides:

5'TATGTGTGGTGGCAGTGGTGGTTCAAATCCACCTGCTCCTACGCCCATT CCAAATG (SEQ ID NO: 36) 3'

3'ACACACCACCACCACCAAGTTTAGGTGGACGAGGATGCGGGTAAGG TTTACGATC (SEQ ID NO: 37) 5'

An NheI-ClaI fragment, containing approximately 1kb of the tbpB gene from pLEM23, prepared as described in Example 7, was ligated to the above oligonucleotides and inserted into pT7-7 cut with NdeI-ClaI, generating pLEM31, which thus contains the 5'-half of tbpB. Oligonucleotides also were used to construct the last approximately 104 bp of the tbpB gene, from the AvaII site to the end of the gene. A BamHI site was incorporated into the 3' end of the oligonucleotides:

5'GTCCAAATGCAAACGAGATGGGCGGGTCATTTACACACAACGCCGATG

ACAGCAAAGCCTCTGTGGTCTTTGGCACAAAAAGACAACAAGAAGTTAAGTAGTA
G (SEQ ID NO: 38) 3'

3'GTTTACGTTTGCTCTACCCGCCCAGTAAATGTGTGTTGCGGCTACTGTC
GTTTCGGAGACACCAGAAACCGTGTTTTTCTGTTGTTCTTCAATTCATCCTAG

(SEQ ID NO: 39) 5'

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A ClaI-AvaII fragment from pLEM23, containing approximately 0.9 kb of the 3'-end of the tbpB gene, was ligated to the AvaII-BamHI oligonucleotides, and inserted into pT7-7 cut with ClaI-BamHI, generating pLEM32. The 1.0 kb NdeI-ClaI insert from pLEM31 and the 1.0 kb ClaI-BamHI insert from pLEM32 were then inserted into pT7-7 cut with NdeI-BamHI, generating pLEM33 which has a full-length tbpB gene under the direction of the T7 promoter.

DNA was purified from pLEM33 and transformed by electroporation into electrocompetent BL21(DE3) cells (Novagen; Madison, WI), to generate strain pLEM33B-1. Strain pLEM33B-1 was grown, and induced using IPTG, as described above in Example 10. Expressed proteins were resolved by SDS-PAGE and transferred to membranes suitable for immunoblotting. Blots were developed using anti-4223 Tbp2 antiserum, diluted 1:4000, as the primary antibody, and rprotein G conjugated with horseradish peroxidase (Zymed) as the secondary antibody. chemiluminescent substrate (Lumiglo; Kirkegaard and Perry Laboratories, Gaithersburg, MD) was used detection. Induced recombinant proteins were visible on the Coomassie blue-stained gels (Fig. 19). 4223 Tbp2 antiserum recognized the recombinant proteins on Western blots.

Example 13

This Example illustrates the generation of an expression plasmid for rTbp2 of *M. catarrhalis* 4223 with a leader sequence.

The construction scheme is shown in Figure 18. Oligonucleotides containing the natural leader sequence of the *M. catarrhalis* 4223 *tbpB* gene were used to construct the first approximately 115 bp of the *tbpB* gene to the *NheI* site. An *NdeI* site was incorporated into the 5' end of the oligonucleotides:

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5'TATGAAACACATTCCTTTAACCACACTGTGTGGGCAATCTCTGCCGTC TTATTAACCGCTTGTGGTGGCAGTGGTGGTTCAAATCCACCTGCTCCTACGCCCAT TCCAAATG (SEQ ID NO: 40) 3'

3'ACTTTGTGTAAGGAAATTGGTGTGACACACCGTTAGAGACGGCAGAA
TAATTGGCGAACACCACCGTCACCACCAAGTTTAGGTGGACGAGGATGCGGGTAAG
GTTTACGATC (SEQ ID NO: 41) 5'

The NdeI-NheI oligonucleotides were ligated to pLEM33 cut with NdeI-NheI, generating pLEM37, which thus contains a full-length 4223 tbpB gene encoding the Tbp2 protein with its leader sequence, driven by the T7 promoter.

DNA from pLEM37 was purified and transformed by electroporation into electrocompetent BL21(DE3) cells (Novagen; Madison, WI), to generate strain pLEM37B-2. pLEM37B-2 was grown, and induced using described above in Example 10. Expressed proteins were resolved by SDS-PAGE and transferred to membranes suitable for immunoblotting. Blots were developed using anti-4223 Tbp2 antiserum, diluted 1:4000, as the primary antibody, and rprotein G conjugated horseradish peroxidase (Zymed) as the secondary A chemiluminescent substrate antibody. (Lumiglo; Kirkegaard and Perry Laboratories, Gaithersburg, MD) was used for detection. Induced recombinant proteins were visible on Coomassie-blue stained gels (Fig. 21). anti-4223 Tbp2 antiserum recognized recombinant proteins on Western blots.

30 Example 14

This Example illustrates the construction of an expression plasmid for rTbp2 of $\it M.$ catarrhalis Q8 without a leader sequence.

The construction scheme for rTbp2 is shown in Figure 20. The 5'-end of the tbpB gene of M. catarrhalis Q8 was PCR amplified from the Cys 1 codon of

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the mature protein through the Bsm I restriction site. An Nde I restriction site was introduced at the 5' end, for later cloning into pT7-7, and the final PCR fragment was 238 bp in length. The PCR primers are indicated below:

Ndel C G G S S G G F N

5' GAATTCCATATG TGT GGT GGG AGC TCT GGT GGT TTC AAT C

3' 5247.RD (SEQ ID No: 42)

5' CCCATGGCAGGTTCTTGAATGCCTGAAACT 3' 5236.RD (SEO ID No: 43).

The Q8 tbpB gene was subcloned in two fragments contained on plasmids SLRD3 and SLRD5, prepared as Example 8. described in Plasmid SLRD3-5 constructed to contain the full-length tbpB gene by digesting SLRD5 with EcoR I and Dra I, which releases the 3'-end of tbpB, and inserting this ~ 619 bp fragment into SLRD3 which had been digested with EcoR I The 1.85 kb Bsm I-BamH I fragment from SLRD and Sma I. 3-5 was ligated with the 238 bp PCR fragment and inserted into pT7-7 that had been digested with Nde I and BamH I, generating plasmid SLRD35B. This plasmid thus contains the full-length tbpB gene without its leader sequence, the direction under of the T7 promoter. DNA from SLRD35B was purified transformed by electroporation into electrocompetent BL21(DE3) cells to generate strain SLRD35BD which was grown and induced using IPTG, as described above in Example 10. Expressed proteins were resolved by SDS-PAGE and the induced Tbp2 protein was clearly visible by Coomassie blue staining (Fig. 19).

Example 15

This Example illustrates the generation of an expression plasmid for rTbp2 of *M. catarrhalis* Q8 with

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a leader sequence.

The construction scheme for the rTbp2 is shown in Figure 20. The 5'-end of the Q8 tbpB gene was PCR amplified from the ATG start codon to the Bsm I restriction site. An Nde I site was engineered at the 5'-end, to facilitate cloning into the pT7-7 expression vector, and the final PCR fragment was 295 bp. The PCR primers are indicated below:

Nde I K H I P L T

5' GAATTCCATATG AAA CAC ATT CCT TTA ACC 3' 5235.RD

(SEQ ID No: 44)

5' CCCATGGCAGGTTCTTGAATGCCTGAAACT 3' 5236.RD (SEQ ID No: 43).

SLRD3-5 (Example 14) was digested with Bsm I and BamH I, generating a 1.85 kb fragment, which was ligated with the 295bp PCR fragment and ligated into pT7-7 that had been digested with Nde I and BamH I. The resulting plasmid SLRD35A thus contains the full-length Q8 tbpB gene with its endogenous leader sequence under the control of the T7 promoter. DNA from SLRD35A was purified and transformed by electroporation into electrocompetent BL21(DE3) cells to generate strain SLRD35AD which was grown and induced using IPTG, as described above in Example 10. Expressed proteins were resolved by SDS-PAGE and the induced Tbp2 protein was clearly visible by Coomassie blue staining (Fig. 19).

30 Example 16

This Example illustrates the extraction and purification of rTbp2 of *M. catarrhalis* 4223 and Q8 from *E. coli*.

pLEM37B (4223) and SLRD35AD (Q8) transformants

were grown to produce Tbp2 in inclusion bodies and then
the Tbp2 was purified according to the scheme in Figure

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22. E. coli cells from a 500 mL culture, were resuspended in 50 mL of 50 mM Tris-HCl, pH 8.0 containing 5 mM AEBSF (protease inhibitor), and disrupted by sonication (3 x 10 min, 70% duty circle). The extract was centrifuged at 20,000 x g for 30 min and the resultant supernatant which contained > 95% of the soluble proteins from $E.\ coli$ was discarded.

The remaining pellet (PPT₁) was further extracted in 50 mL of 50 mM Tris, pH 8.0 containing 0.5% Triton X-100 and 10 mM EDTA. The mixture was stirred at 4° C for at least 2 hours and then centrifuged at 20,000 x g for 30 min and the supernatant containing residual soluble proteins and the majority of the membrane proteins was discarded.

The resultant pellet (PPT₂) obtained after above extraction contained the inclusion bodies. Tbp2 protein was solubilized in 50 mM Tris, pH 8.0, containing 6 M guanidine and 5 mΜ DTT. centrifugation, the resultant supernatant was further purified on a Superdex 200 gel filtration column equilibrated in 50 mM Tris, pH 8.0, containing 2 M guanidine and 5 mM DTT. The fractions were analyzed by SDS-PAGE and those containing purified Tbp2 pooled. Triton X-100 was added to the pooled Tbp2 fraction to a final concentration of 0.1%. The fraction was then dialyzed overnight at 4°C against PBS, and then centrifuged at 20,000 x g for 30 min. The protein remained soluble under these conditions purified Tbp2 was stored at -20°C. Figure 22 shows the SDS PAGE analysis of fractions of the purification process for rTbp2 from strain 4223 (Panel A) and strain Q8 (Panel B). The rTbp2 was at least 70% pure.

Groups of five BALB/c mice were injected three times subcutaneously (s.c.) on days 1, 29 and 43 with purified rTbp2 (0.3 mg to 10 mg) from M. catarrhalis strains 4223 and Q8 in the presence or absence of AlpO₄

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(1.5 mg per dose). Blood samples were taken on days 14, 28, 42 and 56 for analysing the anti-rTbp2 antibody titers by EIAs.

Groups of two rabbits and two guinea pigs (Charles River, Quebec) were immunized intramuscularly (i.m.) on day 1 with a 5 mg dose of purified rTbp2 protein emulsified in complete Freund's adjuvant (CFA). Animals were boosted on days 14 and 29 with the same dose of protein emulsified in incomplete Freund's adjuvant (IFA). Blood samples were taken on day 42 for analysing anti-rTbp2 antibody titers and bactericidal activity. Table 2 below shows the bactericidal activity of antibodies raised to the recombinant tranferrin binding proteins rTbp1 (4223), rTbp2 (4223) and rTbp2 (Q8), prepared as described in these Examples, against M. catarrhalis strains 4223 and Q8.

Example 17

This Example illustrates the binding of Tbp2 to human transferrin in vitro.

20 Transferrin-binding activity of Tbp2 was assessed according to the procedures of Schryvers and Lee (ref. with modifications. Briefly, purified rTbp2 was subjected to discontinuous electrophoresis 12.5% SDS-PAGE gels. The proteins 25 electrophoretically transferred to PVDF membrane and incubated with horseradish peroxidase-conjugated human transferrin (HRP-human transferrin. 1:50 dilution) (Jackson ImmunoResearch Labs Inc., Mississauga, Ontario) at 4°C for overnight. LumiGLO substrate (Kirkegaard & Perry Laboratories, Inc., Gaithersburg, 30 MD) was used for chemiluminescent detection of HRP activity according to the manufacturer's instructions. Both 4223 rTbp2 and Q8 rTbp2 bind to human transferrin under these conditions, as shown in Figure 24.

35 Example 18

This Example illustrates antigenic conservation of

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Tbp2 amongst M. catarrhalis strains.

Whole cell lysates of *M. catarrhalis* strains and *E. coli* strains expressing recombinant Tbp2 proteins were separated by SDS-PAGE and electrophoretically transferred to PVDF membrane. Guinea pig anti-4223 rTbp2 or anti-Q8 rTbp2 antisera were used as first antibody and alkaline phosphatase conjugated goat anti-guinea pig antibody was used as second antibody to detect Tbp2. *M. catarrhalis* strains 3, 56, 135, 585, 4223, 5191, 8185 and ATCC 25240 were tested and all showed specific reactivity with anti-4223 rTbp2 or anti-Q8 rTbp2 antibody (Figure 25).

Table 3 illustrates the ability of anti-rTbp2 antibodies from one M. catarrhalis strain to recognize native or recombinant protein from a homologous or heterologous M. catarrhalis strain.

Example 19

This Example illustrates PCR amplification of the tbpB gene from M. catarrhalis strain R1 and characterization of the amplified R1 tbpB gene.

Chromosomal DNA from *M. catarrhalis* strain R1 was prepared using standard techniques. The design of the oligonucleotide sense primer was based on a region approximately 274 bases upstream of the *M. catarrhalis* 4223 tbpB gene, and the antisense primer was based upon a region approximately 11 bases downstream of the end of 4223 tbpB. The following primers were used:

sense primer (4940): 5' GATATAAGCACGCCCTACTT 3'

(SEQ ID No: 48)

antisense primer (4967): 5' CCCATCAGCCAAACAACATTGTGT 3'

(SEQ ID No: 49).

Each reaction tube contained 10 mM Tris-HCl (pH 8.85), 25 mM KCl, 5 mM (NH₄)₂SO₄, 2 mM MgSO₄, 800 mM dNTPs, 1.0 mg each of primers 4940 and 4967, 10 ng of R1 DNA, and 2.5 U Pwo DNA polymerase (Boehringer

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Mannheim) in а total volume of 100 μl. The thermocycler was programmed for 5 min at 95°C, followed by 25 cycles of 95°C for 30 sec, 50°C for 45 sec, and 2 and a 10 min final elongation min, The amplified product was purified elongation at 72°C. using a Geneclean (BIO 101) according manufacturer's instructions, and sequenced.

A partial restriction map of M. catarrhalis strain R1 tbpB prepared as just described is shown in Figure The nucleotide and deduced amino acid sequences of the PCR amplified R1 tbpB gene are shown in Figure 27. The R1 tbpB gene encodes a 714 amino acid protein of molecular weight 76.8 kDa. The leader sequence of the R1 Tbp2 protein is identical to that of the 4223 and Q8Tbp2 proteins. When the deduced R1 Tbp2 sequence was aligned with the 4223 Tbp2 sequence, it was found to be identical and 88% homologous (Fig. 28). conserved LEGGFYG (SEQ ID No: 50) epitope was present, as found in Tbp2 from other M. catarrhalis strains as well as the H. influenzae and N. meningitidis Tbp2 proteins.

SUMMARY OF THE DISCLOSURE

summary of this disclosure, the invention provides purified and isolated DNA molecules 25 containing transferrin receptor genes of Moraxella catarrhalis, the sequences of these transferrin receptor genes, and the derived amino acid sequences thereof. The genes and DNA sequences are useful for diagnosis, 30 immunization, and the generation of diagnostic and immunological reagents. Immunogenic compositions, including vaccines, based upon expressed recombinant Tbp1 and/or Tbp2, portions thereof, or analogs thereof, can be prepared for prevention of diseases caused by Moraxella. Modifications are possible within the scope 35 of this invention.

TABLE!

BACTERICIDAL ANTIBODY TITRES FOR M. CATARRHALIS ANTIGENS

ANTIGEN'	SOURCE OF ANTISERA 2	BACTERICIDAL TITRE ³ RH408 ⁴		BACTERICIDAL TITRE Q8 ⁵	
	·	Pre-Immune	Post-Immune	Pre-Immune	Post-Immune
TBP1	GP	< 3.0	4.2-6.9	< 3.0	4.46.2
TBP2	GP	< 3.0	12.0-13.6	< 3.0	< 3.0-4.0

- 1 antigens isolated from M. catarrhalis 4223
- 2 GP = guinea pig
- 3 bactericidal titres: expressed in log₂ as the dilution of antiserum capable of killing 50% of cells
- 4 M. catarrhalis RH408 is a non-clumping derivative of M. catarrhalis 4223
- 5 M. catarrhalis Q8 is a clinical isolate which displays a non-clumping phenotype

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TABLE 2

Antigen	Bactericidal titre -	RH408	Bactericidal titre - Q8		
	pre-immune	post-immune	pre-immune	post-immune	
rTbp1 (4223)	< 3.0	< 3.0	< 3.0	< 3.0	
rTbp2 (4223)	< 3.0	10-15	< 3.0	< 3.0	
rTbp2 (Q8)	NT	NT	< 3.0	5.5-7.5	

Antibody titres are expressed in \log_2 as the dilution of antiserum capable of killing 50% of cells

NT = not tested

TABLE 3

ELISA titres for anti-rTbp2 antibodies recognizing native or rTbp2 from strain 4223 or rTbp2 from strain Q8

	Anti-rTbp2 (4223) Antibody Titres		Anti-rTbp2 (Q8) Antibody Titres	
Coated antigen	Rabbit Guinea pig antisera antisera		Rabbit antisera Guinea antise	
Native Tbp2	409,600	1,638,400	25,600	51,200
	204,800	1,638,400	25,600	102,400
(4223)	409,600	1,638,400	102,400	204,800
rTbp2 (4223)	409,600	1,638,400	102,400	204,800
rTbp2 (Q8)	409,600	1,638,400	1,638,400	1,638,400
	102,400	1,638,400	409,600	1,638,400

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CLAIMS

What we claim is:

- 1. A purified and isolated nucleic acid molecule encoding a transferrin receptor protein of a strain of *Moraxella* or a fragment or an analog of the transferrin receptor protein.
- 2. The nucleic acid molecule of claim 1 wherein the transferrin receptor protein is the transferrin receptor binding protein 1 (Tbp1) of the *Moraxella* strain.
- 3. The nucleic acid molecule of claim 2 wherein the transferrin receptor protein is the transferrin receptor binding protein 2 (Tbp2) of the *Moraxella* strain.
- 4. The nucleic acid molecule of claim 1 wherein the strain of *Moraxella* is a strain of *Moraxella* catarrhalis.
- 5. The nucleic acid molecule of claim 4 wherein the strain of *Moraxella catarrhalis* is *Moraxella catarrhalis* 4223, Q8 or R1.
- 6. A purified and isolated nucleic acid molecule having a DNA sequence selected from the group consisting of:
- (a) a DNA sequence as set out in Figure 5, 6, 10, 11 or 27 (SEQ ID Nos: 1, 2, 3, 4, 5, 6, 7, 8, 45 or 46) or the complementary DNA sequence thereto;
- (b) a DNA sequence encoding an amino acid sequence as set out in Figure 5, 6, 10, 11 or 27 (SEQ ID Nos: 9, 10, 11, 12, 13, 14, 15, 16 or 47) or the complementary DNA sequence thereto; and
- (c) a DNA sequence which hybridizes under stringent conditions to any one of the DNA sequences defined in (a) or (b).
- 7. The nucleic acid molecule of claim 6, wherein the DNA sequence defined in (c) has at least about 90% sequence identity with any one of the DNA sequences

defined in (a) or (b).

- 8. The nucleic acid molecule of claim 6 wherein the DNA sequence defined in (c) is that encoding the equivalent transferrin receptor protein from another strain of *Moraxella*.
- 9. A vector adapted for transformation of a host comprising the nucleic acid molecule of claim 1 or 6.
- 10. The vector of claim 9 encoding a fragment of a transferrin receptor protein and having the characteristics of a plasmid selected from the group consisting of pLEM3, pLEM25, pLEM23, DS-1698-1-1, DS-1754-1, pSLRD2, pSLRD3, pSLRD4 and pSLRD5.
- 11. The vector of claim 9 further comprising expression means operatively coupled to the nucleic acid molecule for expression by the host of said transferrin receptor protein of a strain of *Moraxella* or the fragment or the analog of the transferrin receptor protein.
- 12. The vector of claim 11 having the characteristics of plasmid pLEM-29, pLEM-33, pLEM-37, SLRD35-A and SLRD35-B.
- 13. A transformed host containing an expression vector as claimed in claim 11.
- 14. A method of forming a substantially pure recombinant transferrin receptor protein of a strain of *Moraxella*, which comprises:

growing the transformed host of claim 13 to express a transferrin receptor protein as inclusion bodies,

purifying the inclusion bodies free from cellular material and soluble proteins,

solubilizing transferrin receptor protein from the purified inclusion bodies, and

purifying the transferrin receptor protein free

from other solubilized materials.

- 15. The method of claim 14 wherein said transferrin receptor protein comprises Tbpl alone, Tbp2 alone or a mixture of Tbpl and Tbp2.
- 16. The method of claim 15 wherein said transferrin receptor protein is at least about 70% pure.
- 17. The method of claim 16 wherein said transferrin receptor protein is at least about 90% pure.
- 18. A recombinant transferrin receptor protein or fragment or analog thereof producible by the transformed host of claim 12.
- 19. The protein of claim 18 which is transferrin receptor binding protein 1 (Tbp1) of the *Moraxella* strain devoid of other proteins of the *Moraxella* strain.
- 20. The protein of claim 18 which is transferrin receptor binding protein 2 (Tbp2) of the *Moraxella* strain devoid of other proteins of the *Moraxella* strain.
- 21. The protein of claim 18 wherein the strain of Moraxella is a strain of Moraxella catarrhalis.
- 22. An immunogenic composition, comprising at least one active component selected from the group consisting of:
- (A) a purified and isolated nucleic acid molecule encoding a transferrin receptor protein of a strain of *Moraxella* or a fragment or an analog of the transferrin receptor protein;
- (B) a purified and isolated nucleic acid molecule having a DNA sequence selected from the group consisting of:
 - a) a DNA sequence as set out in Figure 5, 6, 10, 11 or 27 (SEQ ID Nos: 1, 2, 3, 4, 5, 6, 7, 8, 45 or 46) or the complementary DNA sequence thereto;

- (b) a DNA sequence encoding an amino acid sequence as set out in Figure 5, 6, 10, 11 or 27 (SEQ ID Nos: 9, 10, 11, 12, 13, 14, 15, 16 or 47) or the complementary DNA sequence thereto; and
- (c) a DNA sequence which hybridizes under stringent conditions to any one of the DNA sequences defined in (a) or (b); or
- (C) a recombinant transferrin receptor protein fragment or analog thereof producible by transformed host containing an expression vector comprising a nucleic acid molecule as defined in (A) or and expression means operatively coupled to the nucleic acid molecule for expression by the host of the recombinant transferrin receptor protein or fragment or analog thereof;

and a pharmaceutically acceptable carrier therefor, said at least one active component producing an immune response when administered to a host.

- 23. A method for generating an immune response in a host, comprising administering to the host an immunoeffective amount of the immunogenic composition of claim 22.
- 24. A method of determining the presence, in a sample, of nucleic acid encoding a transferrin receptor protein of a strain of *Moraxella*, comprising the steps of:
- (a) contacting the sample with the nucleic acid molecule of claim 1 or 6 to produce duplexes comprising the nucleic acid molecule and any said nucleic acid molecule encoding the transferrin receptor protein of a strain of *Moraxella* present in the sample and specifically hybridizable therewith; and
 - (b) determining production of the duplexes.

- 25. A diagnostic kit for determining the presence, in a sample, of nucleic acid encoding a transferrin receptor protein of a strain of *Moraxella*, comprising:
 - (a) the nucleic acid molecule of claim 1 or 6;
- (b) means for contacting the nucleic acid molecule with the sample to produce duplexes comprising the nucleic acid molecule and any said nucleic acid present in the sample and hybridizable with the nucleic acid molecule; and
- (c) means for determining production of the duplexes.

AMINO ACID SEQUENCES OF A CONSERVED PORTION OF Tbp1 PROTEIN FOR CONSTRUCTION OF DEGENERATE PRIMERS USED IN PCR AMPLIFICATION OF A PORTION OF THE M. cattarhalis 4223 tbpA GENE.

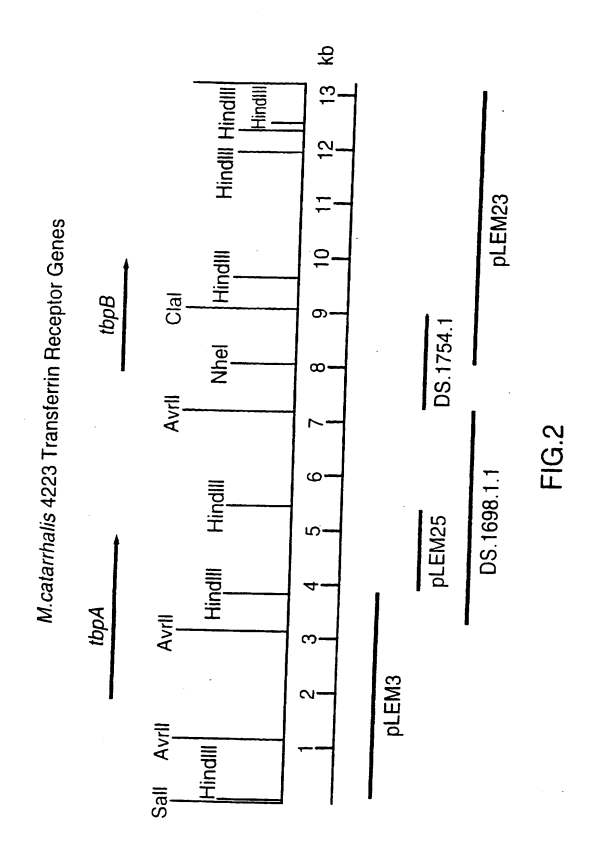
NEVTGLG

SEQ ID NO: 17

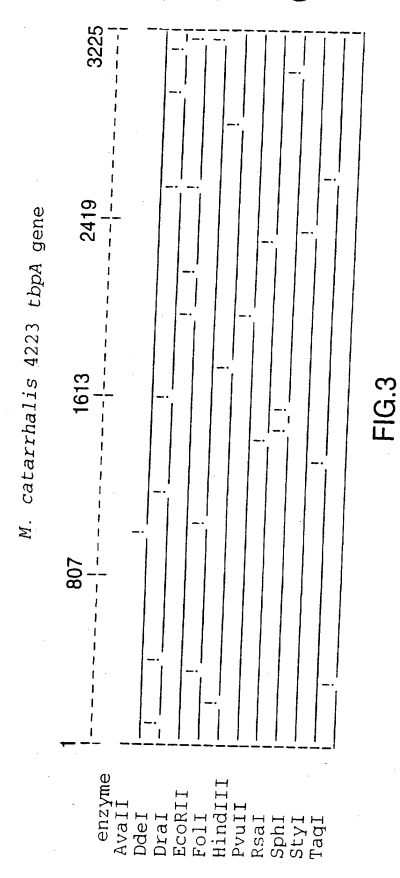
GAINEIE

SEQ ID NO: 18

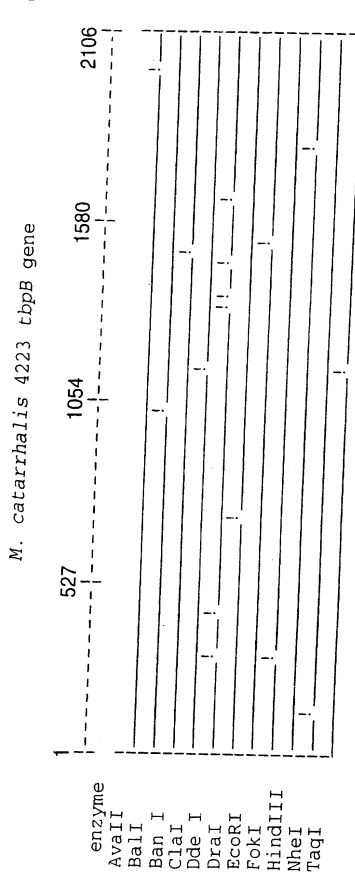
FIG.1



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Leu

CTGLeu

162 TTG

Leu Val

Val

Val

CTTLeu

AAC Asn

Thr

Lys

ACA

216 **ACA**

GAA Glu

AAC

ညည Ala

AAA Lys

Val

Asn

tbpA gene catarrhalis 4223 Sequence of M.

TATTTTGGTAAACAATTAAGTTCTTAAAAACGATACACGCTCATAAACAGATGGTTTTTTGGCATCTGCAAT

TTGATGCCTGCCTTGTGATTGGGTTGTGTGTGTGTGTTCAAAGTGCAAAAGCCAACAGGTGGTCATTG

TCCTCC AAA AAA LysAAA Gln Asn Asn TCA AAA CAA AAC AAC

Lys

Ser

CAA Gln

AAT Asn

MET

CAA GTA TTA Gln Val AAA LysSer Ser Lys Lys

81

Ala GTG Val CAG Gln ACG Thr ATC Ile AAC Asn CTT Leu

Leu

GGTGly

TTGLeu

TCT Ser

Leu

Ser

Leu

225

AGT

AAG

GCA

CCG

AAG Lys

GAT

ACG Thr

Asp

GAT ACA Thr GAG

Asp Ala Glu Ala

GTA Val GTT Val

Thr

GAA

CGTArg CCAAC Asn AAA Lys 189 AAG Lys CCG Ala ACA Thr

4	

Pro

Leu Gln

	•
270	324
CTA	GGT
Leu	G1v
GTG	CAA
Val	Gln
CAA	GAG
Gln	Glu
GAA	GTT
Glu	Val
aaa	GTG
Lys	Val
AAT	GCT
Asn	Ala
ATC	ATT
Ile	Ile
ACC	GGC
Thr	Gly
GAG	CCT
Glu	Pro
243	297
GCC	GAC
Ala	Asp
ACT	tat
Thr	Tyr
AAA	cgc
Lys	Arg
GTC Val	ACA Thr
Grc	TTA
Val	Leu
AAG	GAC
Lys	Asp
GGT 1	CGA Arg
Crr	ATT
Leu	Ile
GGG	AAC Asn
	SUBSTI

378	432
GCG	GTG
Ala	Val
GTG Val	CCT
CGT (GGC G1v
aat Asn	432 CAA GGC CCT GTG Gln Glv Pro Val
	CTA
GAT AAA)
Asp Lys	(1)
CGT GGT ATG Arg Gly MET	G CAC TAT G
GGT	CAC
G1y	His
cgt	cAG
Arg	Gln
351 ATT Ile	405 A GCC CAG (1)
TCT	c AAT CAA
Ser	e Asn Gln
GGC TAT $G1y Tyr$	AAT Asn
GGC	ATC
G1y	I1e
TCA Ser	${\color{red}GGC} \\ {\color{blue}G1} \\ {\color{blue}y} \\$
AGC	GAT
Ser	Asp
GCA	GTT
Ala	Val
GGG	TTG
G1y	Leu
CGT	GTA
Arg	Val
TUTE SHEET	(DIII E 26)

486 AAT Asn GAA $\mathbf{TAC} \\ \mathbf{TYr} \\$ **GAA** Glu ATA Ile **GAA** Glu **AAC** Asn 459 **GGG** Gly TyrTAT **AAT** Asn AAA Lys **GGC** Gly

540 GGG G1y TAC Tyr TCA AAT Asn 513 GGT G1y AAA Lys GAG CGC

810 CGA Arg

> AAC Asn

AAT Asn

GAC Asp

ACT Thr

ACC Thr

GTG Val

783 GCG Ala

> AGA Arg

GAT Asp

AGT Ser

CAA Gln

AGC Ser

 ${\rm GGT} \\ {\rm G1y}$

CAG Gln

FIG.5C

₩ 4 10	~ • • • •		
594	648	702	756
AAA	AAC	CTT	TAT
Lys	Asn	Leu	TYr
GCC GAT GAC ATC ATC	A AAT A	GGT	3CC
Ala Asp Asp Ile Ile		G1y	Ala
ATC	AAA	AGC	GAT GAT (
Ile	Lys		Asp Asp ?
GAC	AGT	TTT Phe	GAT
GAT	GCC AGT A	TCT Ser	CAT
	\mathtt{TAT}	GGT	GCA Ala
ACC	GCC	GCA	AAG
Thr		Ala	Lys
AAA	ACC	AAG	TAC
Lys		Lys	Tyr
ACC	AAA	GGC	GAA
	Lys	G1y	Glu
567	621	675	729
GTT	ACC	GCA	CAA (
Val	Thr	Ala	Gln (
TTT	CAG	GCA GCA ($_{\rm G1Y}$
Phe	Gln	Ala Ala A	
GCA	GTG	GCA	GT
Ala	Val	Ala	
GTG	GGC (Gly)	GTG	CGC
Val		Val	Arg
TCT	TGG	TCT	GAC
Ser	Trp	Ser	Asp
TCT GGC TCT G	GAT	GTT AAT TCT GTG (ACC GAC CGC C
Ser Gly Ser V		Val Asn Ser Val <i>i</i>	Thr Asp Arg A
TCT	AAA	GTT	TAC
Ser	Lys	Val	Tyr
TTA	GAT GGT AAA GAT TGG	lgg	ATC
	Asp Gly Lys Asp Trp	lrp	Ile
GCA	GAT	GCA 7	ATC
Ala	Asp		Ile

864 GCT Ala 918 AAT Asn GCG Ala GAT Asp GAG AAT Asn AAT Asn ${\tt GGT} \\ {\tt G1y}$ AAT Asn 837 GCC Ala TGT GAA Glu CAA Gln AAT Asn TTA Leu

AAC

FIG.5D

972 GAC Asp CAA Gln ACC Thr CTC Leu CCA Pro AAC Asn CCA Pro CTTLeu Arg 945 CGC Asn AAC CCT Pro $\text{GGT}\\\text{Gl}\gamma$ ACA Thr TAT Asp GAT AAA Lys GTC Val

1026 TATTyrCAC His AAG Lys GAT Asp AAC Asn CTA Leu CAG Gln TAT 999 GGT G1y CCA Pro CGC Arg CTT CTG TTA Leu TCC Ser AAA Lys AGC

1080 ACC Thr AAA Lys Asp GATCAA Gln ATG MET GCC Ala TAC Tyr AAC Asn 1053 CAA Gln Lys AAA ACC Thr ATC Ile GAA TAT Tyr GTG Val GGT GGT

Asn AGC Ser CTC AGG Arg TCA Ser AAA Lys GAA Glu 1107 GAC ATT Asp Ile CAT His GTT Val ACG Thr

CTG

TAT Tyr

GCT Ala

CCT

GAT ACC Asp Thr 1188 ATT Ile CGC GAA Glu ${\tt GGT} \\ {\tt Gly}$ CTT Leu AAT Asn GGC AAT Gly Asn 1161 CAA Gln TAT TyrTAT GGC Gly AAT Asn CAA Gln

1242 GAT GT'A Val GGC Gly CAT His GCT TAT Tyr AAC Asn 1215 ATC . Ile A GGC Gly TAT GGT TCA Ser

FIG.5E

296 GGT Lys Ser Asp GAC TyrTAT GTT Val TAT Γyr GAA Glu Leu 999 Leu Arg CGC GAC Asp AAA Lys CAA Gln CAC His AAA Lys3AA

1350 Asp GAC Gln CAA AAG LysAsp TAT Tyr TCT Ser Val CGTVal Asp Asp GAT Phe TTT Trp TGG Lys Asn AAT

Lys Asp GAC CAC His Pro SSS TyrTAT ACC TCA CysCAC His 1377 Thr AAC Asn ACC Thr CTG CAG Gln Ser Arg CGC Leu

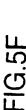
Asn Asn Asp Val GTG GAG Glu AAA Lys GTA Val TCG Ser Phe CCTPro1431 AAA Lys Asn AAT GTC Val Asp GAT Pro ACG Thr TGT AAT Asn

MET AAA Lys AAA Lys AAC Asn TTT Phe GTC Val Ala AAA Lys ATC 1485 Leu CAC His CAG Gln Glu AAA Lys PAC Pyr

AAT Asn AAA Lys GAT Asp TAT G1yCGC GTT Val CAA Gln CTG Leu .539 AAC Asn Ile His CAT His ACG Thr AGT Ser GGC Gly Leu

620 AAA CAA TAT TCT Ser CAT His ACC Thr TTG Leu CGT Arg 593 TAT Tyr Asp GAA Glu Arg CGT AGC Ser CTG Leu AGC Ser Ser





CCC Pro AAG Lys Phe TTTAAG Lys GAT Asp CCA Pro TTG Leu AAC Asn AGT Ser CCA Pro CCA Pro ACC Thr TAC GAT Asp CTT Leu

GAC Asp CAT His GGT Gly TAT GGT Gly TAT Tyr GCT GAT Asp 1701 CTT Leu TGC Cys ATT Ile CCC Pro AAA Lys AAC Asn AAC Asn TCA

AAA AAA Lys Lys ATC Ile GCC TTT Phe AAT Asn CAA Gln TAT ACT Thr 1755 AGC Ser AAC Asn AAA Lys GCC Ala AAC Asn TGT GCT CAG Gln CCA

CCGln Ala CAA TAT Tyr GAT Asp ATT Ile AAG Lys GAT Asp ACC Thr AAT Asn 1809 ACC Thr AAA . Lys ' CAA Gln AAC Asn TAC Tyr CAA Gln GAG ATA Ile

1890 GAG Phe Pro CCCAAA Lys CTA Leu AÇC Thr AGC Ser Asn AAC CCC 1863 CAA AAC Gln Asn AAA Lys GAT Asp TAT Tyr CAA Gln GAC

2991944 Leu GAA Glu GAC ATA Ile AAG Lys AAC Asn TAC AAA Lys GAA Glu Gln CAA GGG G1y TTG AGT Ser CAA Gln AAA Lys Lys

1998 AAC Asn





Asp

Tyr

Lys

Leu

Asp GAC Asn ACT Thr TGGTrp $\begin{array}{c} \text{GGT} \\ \text{G1} \end{array}$ CCC Ala $_{
m LCG}$ Trp Glu GAA Asn TTA Leu Asp GAT Lys TYYTATGCT AAA Lys Phe

Asn Pro CCA CAG Gln TAT TyrATC Ile AAAT Asn GAT Asp ACG Thr 2025 Lys AAA AAT Asn Asn AAT Gln CAA Ser

2052 GCA

2106 Asp GAT Ala $\Gamma Y \Gamma$ AGC Ser AAC Asn ACC Thr GAG Glu AGC TAT TyrAAA Lys 2079 $\mathbb{L}G\mathbb{L}$ AAA Lys GAC Asp Asp GAT Lys AAA GTC GTG Val Thr

2160 GAC AAA TTC Phe TATTYrAAT Asn GAT Asp 2133 GGTG1ySer CAC His CGC Arg Thr ACC Thr TCA $_{\rm LGC}$

GAC TAT GCT GGTG1yCTG Len GGG G1yLeu Asp GAT 2187 Val TATTyrAAA Lys AAT Asn ACC ATG MET AAC Asn

2268 CAG AAC Asn AGC Ser AGT Ser AAC Asn GAC Asp GTA Val 2241 TTG Leu GTG Val GAT Asp Ser AAA Lys AAA

FIG.51

GAC Asp CTG Leu Trp 16GAsn AAT ACC Thr CCC Pro 2295 AAG Val GTC Val GTG Val GGC Gly TTT Phe AAT Asn TGGTrp TCT Ser

CGC GGC Gly TAT Tyr ATG MET Glu GAA TCT Phe $\Gamma\Gamma\Gamma$ AGT Ser CCA 2349 Pro ATG Arg CGC Phe TTTGGC Gly CAA Gln TCG AGC Ser

TAC Tyr Leu GGT Gly AAG Lys TGT Cys GGC Gly CAT His CAA Gln 2403 ACG Thr GGC Gly AAA Lys GGT Gly ATC Ile ACC Thr GTA Val GGC Gly

2484 AAC Asn TCC AAA Lys GAA Glu Pro CCT AAA Lys CTA Leu AAG Lys ACC 2457 Thr CAA Gln CAT His GTC ACT Thr CAG Gln CAG Gln TGT Cys

2538 TAT Ser GTT Val GAG Glu CTT Leu AGT Ser GGC Gly Leu TTA CAC His 2511 CAT AAC Asn Leu TTA ACT Thr GCG Ala GGA Gly ATC Ile GAA Glu CAA Gln

2592 AGA ATT Ile GAG Glu GAA Glu AGT Ser AAA Lys GGTGly GTT Val 565 ATT Ile Leu Asp GAT ACC Thr TATTyrCGC Arg AAT Asn AAA Lys Phe

646 TTT Leu GAT Asp GGTGly AAA Lys GGT Gly CGT Arg CAG Gln AAA Lys2619 CCC G1yAAT Asn GGT CAA Gln ACC Thr Leu

AGA

GGC

CTT

Leu



Asn AAC ATT Ile GGC Gly ren , Asp GAT GAT Asp CAA Gln GGA Gly AAT Asn

Leu ACA Thr TCA TAC TTA Leu GGA Gly TyrLeu Arg AGT Ser AAT Asn GTC Asn Leu

808 AAC Asn GGA Gly GCA Ala Leu ACT Thr CCA Pro Asn AAC TTA 2781 ACC Lys AAA GGA Gly AAA Lys GTT Val GAT Asp GTT Val AAA Lys AAC Asn

GAT TAT G1yCTT G1yGTG Val GTG Val TyrTAT Arg CGT 2835 . TCT Ser Pro GCC GAT Asp Phe Leu

AAA GAT Asp TCT CAT His ACC Thr Phe GCC 2889 AAC Asn TGG Trp AAA Lys CAA Gln AGC Ser CCA

AAC Asn AAT Asn GGT Gly Leu Asn AAC AAG Lys LLC Leu CTT GAG Glu AGC Ser Pro Asn

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FIG.5.

3024 GGT G1y Ser TTG GAT Asp CTT ACA Thr CAA Gln TGG Trp CCG 2997 ACG Thr TCC Ser AAA Lys GCA Ala AAA Lys ACC Thr GCC Ala CAA Gln AAA Lys

3078 Val AAT Asn T'AC T'yr GTG Val GGC Gly GCT CGT Arg T'IG Leu 3051 ACC Thr TTT Phe AAT Asn GAT AAA Lys ATA Ile AAC Asn GTA Val TAT Tyr

Val GGG Gly GAA Glu GCA Ala ACA Thr CAA Gln CGC Arg TTA Leu 3105 GAG GCT Glu Ala TGG Trp ACT Thr ACC Thr TAC Tyr TAT ACC Thr AAT Asn

GCC TAT Tyr Arg GGT Gly TAT CAT His AAG Lys GATAsp 3159 CAA AGC Ser CTG Leu GGA Gly ACA Thr CAT His CAG Gln AATASn

TAA AAG Lys ATG MET Leu GCA Ala TTG Leu CAA Gln TAC AAT Asn

GGA Gly



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270 GAT Asp

Sequence of M. catarrhalis 4223 tbpB gene

TGTCAGCATGCCAAAATAGGCATCAACAGACTTTTTTAGATAATACCATCAACCCATCAGAGGATTATTTT

54 TTA Leu
TA
IC T
C GTC a Val
. GCC
TCT
ATC Ile
CA 1a
GTG G Val A
TE G
7 G TGT u Cys
A CTG TGT GTG of Leu Cys Val
ACA
ACC ACA Thr Thr
TTA ACC Leu Thr
ro I
L B C
C A S
CAC His
AAA Lys
ATG MET

(108	CCA	Pro
		ATT	Ile
			Pro Ile Pro
		ACG	Thr
	1		a Pro Thr
	[ر []. المالية	Ala
	. E	CLA CLI GCI	Pro .
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	GGT	Ser Gly	7 7
	AGT	Ser	1)
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E	.T.55	Gly	1
E	757	Cys	
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ACC	170 OCT	Tur	

162 GAT Asp
ACT Thr
GGT
GGC (
GCT (Ala (
AAT C Asn A
GGT A
ACT G Thr G
AAC A Asn T
135 GGC A Gly A
1 ACT G Thr G
AAT A Asn T
GGT A Gly A
TCA G
GGT T(Gly S(
T AGC a Ser
T GCT n Ala
AAT Asn

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GAA AAA Glu Lys GAG AAA AAT (Glu Lys Asn (ACT Thr CCA Pro GTA Val 243 GAT Asp CAA Gln TAT (CCA AAA ? Pro Lys ' GAG Glu CCA ACA AAC Asn

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AAA	ACC	TCG	GCG	GAT	CAG
Lys	Thr	Ser	Ala	Asp	Gln
AGT	ATT	TTT	GTA	TCC	TTT
Ser	Ile	Phe	Val		Phe
TTG	ATC Ile	CCA '	AAT Asn	ATC Ile	GAG Glu
324 ATG GCT TTG AGT AAA MET Ala Leu Ser Lys	AAT Asn	TTG	ATG	GAA Glu	CAT His
ATG	AAA	CCA	AAA	AAA	AGC
	Lys	Pro	Lys	Lys	Ser
324 TAT GGC ATG GCT TTG AGT AAA Tyr Gly <u>MET Ala Leu Ser Lys</u>	GAA G1u	TCG	GCA Ala	AAT Asn	AAA Lys
TAT	GAT	AAA	ATA	GGT	CGT
Tyr	Asp	Lys	Ile	G1y	
$_{\rm G1\gamma}$	TTA Leu	AAA Lys	TAT ATA GCA AAA ATG AAT GTA GCG Tyr Ile Ala Lys <u>MET Asn Val Ala</u>	AAA Lvs	GTG
ATG MET	CCA	GGT G1y	GGC	AAG Lys	GCT GTG CGT AAA AGC Ala Val Arg Lys Ser
297	351	405	459	513	567
GCC	ACG	GCA GAA GGT AAA AAA TCG CCA TTG	GAT	ATT	GAA
Ala	Thr	Ala Glu Gly Lys Lys Ser Pro Leu	Asp	Ile	Glu
CCT	GAC	GCA	CTT	AGA	AAA
Pro	Asp	Ala	Leu	Arq	Lys
GAA	CAA	GTT	TTG	GAC	ATC
Glu	G1n	Val	Leu	Asp	Ile
CAA	CGA	CAA	AAA	GGT	CAA
	Arq	Gln	Lys	Gly	Gln
ATT	AAC	AAA	GAA AAT AA	ATT	AAA
Ile	Asn	Lys	Glu Asn Ly	Ile	Lys
TCC	CAC	AAA	GAA	GCC	GCC
	His	Lys	Glu	Ala	Ala
TCA	CTA	GGT	GTA	AAT	CTT
Ser		Gly	Val	Asn	Leu
AAA GTT TCA TCC ATT CAA GAA	ATT AAT CTA CAC AAC CGA CAA GAC	GAC	TTA GAT GTA	AAA	GAA
Lys Val Ser Ser Ile Gln Glu	Ile Asn Leu His Asn Arg Gln Asp	Asp	Leu Asp Val	Lys	31u
AAA	ATT AAT CTA CAC AAC CGA CAA GAC ACG CCA TTA GAT GAA AAA AAT ATC ATT	TTA GAC GGT AAA AAA CAA GTT	TTA GAT GTA GAA AAT AAA TTG	GAT AAA AAT GCC ATT GGT GAC AGA ATT AAG AAA GGT AAT AAA GAA ATC	GAA GAA CTT GCC AAA CAA ATC
Lys	Ile Asn Leu His Asn Arg Gln Asp Thr Pro Leu Asp Glu Lys Asn Ile Ile	Leu Asp Gly Lys Lys Gln Val	Leu Asp Val Glu Asn Lys Leu	Asp Lys Asn Ala Ile Gly Asp Arg Ile Lys Lys Gly Asn Lys Glu Ile	Glu Glu Leu Ala Lys Gln Ile

918 GCA Ala

TGG Trp 1

GGC Gly

GCA Ala

CAA Gln

TCT Ser

891 AAC Asn

AAA

AGC GAA GTG Ser Glu Val

AAC CGA TTT

Asn

GAA Glu

648 ACC Thr	702 AAT Asn	756 GTG Val	810 GAT ASP	864 AGA ARG
A ACA A	702 GCG AAT Ala Asn	756 CCT GTG Pro Val	G CCC ACA CAA J Pro Thr Gln	AGA Arg
3G2	TTG	GGC Gly	ACA Thr	AAC Asn
GAC	TAC Tyr	TTA Leu	CCC	GCC Ala
AAT GAC (Asn Asp (TAC Tyr	s AAT TTA o	TTG Leu	GTT Val
TCA Ser	GGT '	TG(Trr	ACC GCC AAA GAG TTG Thr Ala Lys Glu Leu	ATG ACC GAT GTT GCC AAC AGA MET Thr Asp Val Ala Asn Arg
CAT	TAT (Tyr (729 ACA GAC AAA CTT Thr Asp Lys Leu	AAA Lys	ACC Thr
TTT Phe	GAT	AAA Lys	GCC Ala	ATG
ATT TTT (Ile Phe 1	GTT GAT Val Asp	GAC Asp	ACC Thr	TTT Phe
621 GAA AAC AAA A Glu Asn Lys I	675 TAT TYr	729 ACA Thr	783 ACG Thr	837 GAC Asp
AAC Asn	AAA Lys	AAAL ys	ACA Thr	TGG Prp
GAA Glu	TTA Leu	GTC	GGC Gly	GGA CAT Gly His
CTG	GAT Asp	ACC	AAT Asn	GGA Gly
TCA Ser	CGA Arg	CTA Leu	TAT Tyr	AAA Lys
TCA	ACA	TAT TYr	TTT Phe	\mathtt{TAT}
TTA	ACC Thr	AAT Asn	GTG Val	AAA Lys
GTA Val	GCA	GGC Gly	GGT	GTC Val

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AAA Lys

CAA Gln

1242 GAG GAG Glu Glu

TTT Phe

GGT Gly

AAC Asn

GAT Asp

AGT

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972	1026	1080	1134	1188
T GAT	AAG GAA	GGC AAT	CGC TTC	TTT ACC
o Asp	Lys Glu	Gly Asn	Arg Phe	Phe Thr
CC Pr		66	CG	TT
GCC CCT	TTT	AAG	AAC	CCC
Ala Pro	Phe	Lys	Asn	Pro
TCT	AAT	CAT	GGC	CAC
Ser	Asn		Gly	His
GAC	GTT	GC	CAC	AAA
Asp	Val	rg	His	Lys
AAA GAA GAC TCT	ACT	GAC	ATC	AGC
Lys Glu Asp Ser	Thr	Asp	Ile	Ser
AAA	TTT ACT GTT.	CAA	AAT	ACA AGC AAA CAC CCC
Lys	Phe Thr Val	Gln	Asn	Thr Ser Lys His Pro
ACT	3AG	CTA CAA GAC C	GCC AAT ATC CAC GGC	AAT GAC
	31u	Leu Gln Asp A	Ala Asn Ile His Gly	Asn Asp
TTA	AGT (AAC	GAT	AAT
	Ser (Asn	Asp	Asn
945 TTA Leu	999 AGC Ser	1053 TTT AGT Phe Ser	1107 GAC ATC Asp Ile	
CGC Arg	CAT His	TTT Phe	1 GAC Asp	1 AAT Asn
AAC	TAT GGC (CTG	\mathtt{TAT}	AGC
Asn	Tyr Gly 1	Leu		Ser
TAC	\mathtt{TAT}	AAG	CGC	GCA
Tyr		Lys	Arg	Ala
GAA	3AA	GGT	GAA	ACC
Glu	31u	G1y	Glu	Thr
GAT	GGT	ACA	ACC	GCC
Asp	Gly	Thr	Thr	
TCA AAA GAT GAA	AGC GGT (AAA AAA TTA ACA GGT AAG	ACA AAA ACC GAA CGC	AGT GCC ACC GCA AGC AAT AAA
Ser Lys Asp Glu	Ser Gly (Lys Lys Leu Thr Gly Lys	Thr Lys Thr Glu Arg	Ser Ala Thr Ala Ser Asn Lys
TCA	CAT	AAA	ACA	GGC
Ser	His	Lys	Thr	Gly
TCT	GGT	AAA	GTT	CGT
Ser	Gly	Lys	Val	

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1296	1350	1404	1458	1512	1566
GGT GCT	TAT GCA	GAA AAA	GTC ATT	CCA GAG	ATG GTG AAT GAT GAA GTT AGC GTC
Gly Ala	Tyr Ala	Glu Lys	Val Ile	Pro Glu	MET Val Asn Asp Glu Val Ser Val
GG1 G1 ₂		GAZ Glu	GT(Va]	CC	AG(Ser
TTT	GCC	ACC	ACC	AAG	GTT
Phe	Ala	Thr	Thr	Lys	Val
GTC	GAT	TTT	TCT	GAC	GAA
Val		Phe	Ser	Asp	G1u
TTT GGC GTC	TTA (CCA	GGT	AAA	GAT
Phe Gly Val	Leu	Pro	G1y	Lys	Asp
TTT	ATC	ACC	GTC TTA GGT TCT ACC	ACC	AAT
Phe	Ile	Thr	Val Leu Gly Ser Thr	Thr	Asn
CTC	GCC ATC	ACA TTC ACC CCA TTT	GTC	TTC	GTG
	Ala Ile	Thr Phe Thr Pro Phe	Val	Phe	Val
AAA CTC 1 Lys Leu F	GAA Glu	ACA	TTG	AAT GAA TTC ACC AAA GAC AAG Asn Glu Phe Thr Lys Asp Lys	ATG
AAC	ACC	ACC	AAA	AAT	TTG
Asn	Thr		Lys.	Asn	Leu
1269	1323	1377	1431	1485	1539
AAT GAC	GAA AAA	AGT AAC GCA	GCC AAA A	ACC AAA	GAG ACT
Asn Asp	Glu Lys	Ser Asn Ala	Ala Lys I	Thr Lys	Glu Thr
	GAA Glu	AAC Asn	1 GCC Ala	ACC Thr	GAG Glu
ACC	GAG	AGT	AAT	GAT GCC	GGC
Thr	Glu	Ser		Asp Ala	Gly
TTA	GCT	ACA	GGC Gly	GAT	GCG Ala
TTC TTA	AAA	AAT	TTT	ACT	GAA
Phe Leu	Lys	Asn	Phe	Thr	Glu
AAA	AGT	TTT	AAC	CCT	AAC
Lys	Ser		Asn	Pro	Asn
GGT.	GAG AGT A	ACA Thr	GAT AAC Asp Asn I	GTG Val	ACA Thr
GCA Ala	CGA Arg	GGG Gly	CTG Leu		GCC ACA AAC GAA GCG Ala Thr Asn Glu Ala
CTG	AAA	CTT	CAA	GAT TTG	TCT
	Lys	Leu	Gln	Asp Leu	Ser

FIG.6F

Leu CTTGAG GGT Gly TTTPhe CTA AAA Lys Leu 1593 GAA TAC TTTGGC AAA AAC G1yThr

Lys GAG AAA CGC G1yACA Thr ACC Thr GCT Ala ACC Thr CGC Arg GAA Glu CCC GlyCAA Gln Leu Phe Val AGC Ser AGC Ser

GGA GlyGTA Val Trp TGG AAC Asn GGG Gly TTG Leu TAT Tyr GCC AAA Ala Lys Lys ACA Thr GGC Gly ACA Thr ACC Thr CCA Pro GTA Val

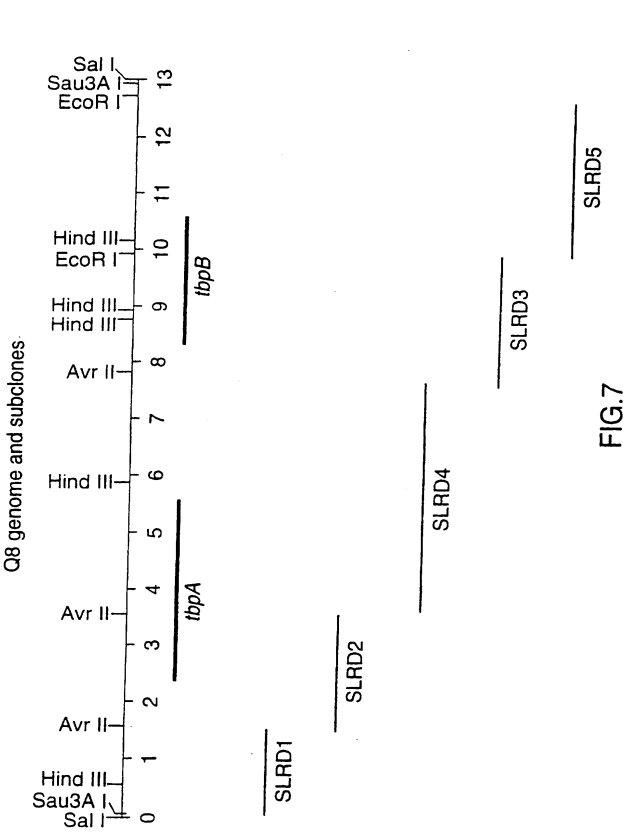
1782 GAT CCC GAT Asp ACC Thr Phe TTTAGC Ser AAA Lys GGA Gly ACA Thr 1755 GGC Gly ACG Thr GGA Gly ACA Thr GAC Asp AAG Lys GGA G1yThr

AAA Lys GGT Gly AGC Ser GTCVal TCA AAA Lys AAT Asn GGA Gly809 Phe GAT Asp ATT Ile GAC Asp Phe GAT Asp Val

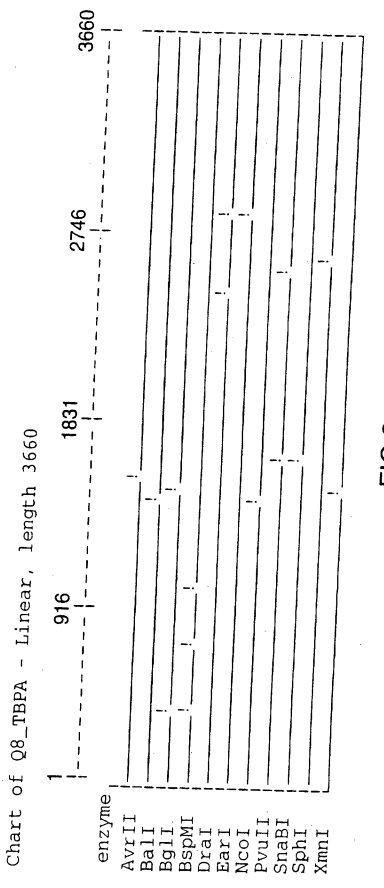
AAT CAA Gln GGT Gly ACA Thr AGC Ser 863 Val Pro GAC Asp CAA Gln

FIG. 6G

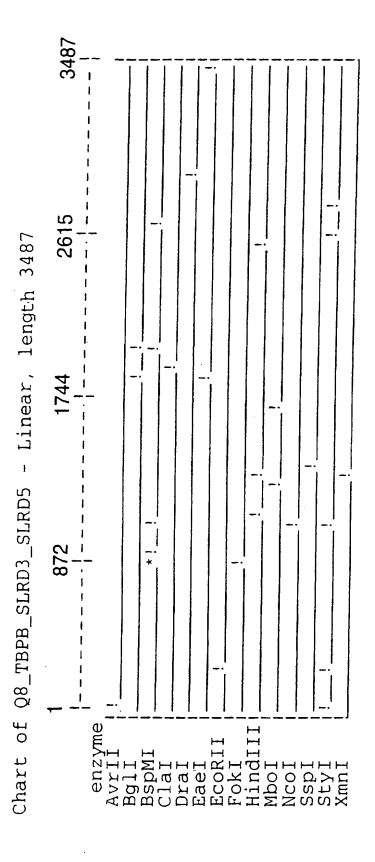
GCG GAC GCA GGA GGC TAC AAG ATA Ala Asp Ala Gly Gly Tyr Lys Ile	ACA GGC AAA TCC ATC GCC ATC AAA GAT GCC AAT GTT ACA GGG Thr Gly Lys Ser Ile Ala Ile Lys Asp Ala Asn Val Thr Gly	GGC GGG TCA TTT ACA CAC AAC GCC Gly Gly Ser Phe Thr His Asn Ala	2106 A AAA AGA CAA CAA GAA GTT AAG r Lys Arg Gln Gln Glu Val Lys
A GC s Al	C AT		ACA Thr
AAA Lys	GCC	ATC	GGC G1y
ACA GCC AGC ACC ACC Thr Ala Ser Thr Thr	1971 A TCC ATC S Ser Ile	2025 GCA AAC GAG ATG Ala Asn Glu MET	2079 GTG GTC TTT Val Val Phe
AG(AAA	GC?	GTG
	Lys	Ala	Val
GCC	660	AAT	Ser V
Ala	G1y	Asn	
ACA	ACA	CCA AAT	AAA GCC
Thr	Thr	Pro Asn	Lys Ala
GGG	AGT	$_{\rm G1Y}^{\rm GGT}$	AAA
Gly	Ser		Lys
ACA GGG	AGC	TAT	AGC
Thr Gly	Ser	Tyr	Ser
TGG	TCT AGC AGT	TTT	GAC
Trp	Ser Ser Ser		Asp
GGC	GAT	GGC	GAT
Gly	Asp	G1y	Asp



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Q8 thpA gene sequence

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FIG. 100

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VAL GLU GIN GLY ARG GLY ALA SER SER GLY
GTTGAGCAGGTCGTGGGCAAGCTCAGGC

ARG GLY MET ASP LYS ASN ARG
ATTCTATTCGTGTATGGATAAAATCGT
610
620

GTGGCGGTATTGGTTGATGGCATCAAT

C A A 660

> ALA GIN HIS TYR ALA LEU GLN GLY PRO VAL CCCAGCACTATGCCCTACAAGGCCCTGTG 670

G A T 840

0000 G G T GCCGCA TAT AAAAAT ASIN 0 0 0 GCA

GTCCGC AAT ASN GAA GLU TAC ø G B Ø AA G A \mathcal{O} ASN Ø \mathcal{O}

T C CSER

G A G A

GCA

A A

园 ALA GLY GAA

TTA GGGGCA TACGGC SER

AAA G

TGGG GAT AAA G AA LYS IE E Ø \mathcal{O} E--GAC

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MAL

AGT TAT C C CZ, A A Ø 880 ں Æ, CAG <u>က</u>

GGTCTT

TATCAGGGT

FIG. 10E

TTT GATGCC LYS AAGGCAGGT Ę--1 GA GCACAT ASN GAA AGAGCGGTGGCAACC TACAAG ALA GGT ALA VAL TYR 133 VAL CGT SER ALA ARG WAL TGG SER ASIN AAT ASIN

ACTGACCCAAATAACCCAAAATTTTAAT.

1060
108

LA ASN GLU CYS ALA ASN GLY ASN TYR GLU C'A A A T G A T G C C A A T G G T A A T T A T G A G 1090 1110

AAC

3 G G T C A A A 1130 GIN GCGTGTGCTGCC GLY ALA

AATGTGCGTGATAAG WAL 混 A C AAGCCA PR PR CAA(

LYS VAL ASIN

AAC ACA GTCAATGTCAAAGAT 1180

PR0

ASIN

PR 084

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ARG

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TCCCAAACCCACTCACCCAAGAC 1210 1220 GLN 强 E

LEG AGCAAATCCTTA E SER LYS

GGT CTTCGC

TAT

PR PR

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AAGCACTATGTCGGTGGT

AACGAT

CAG

ASIN

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SS SS GIN GAAATC GLU

ACCAAACAA 1310

FIG. 100

AAATCA CTTGGTGAA LYS Ø 园 AATAAC ASN C T G 1350 TAT 1410 ASN CAAGATAAAACCGTGCCTGCTTAT ACGGTT GGCTAT CAAGGC M TYR GLY GIN GLY CAAGCCAAT ASIN AACCATGGC ASP GIN ASIN 7 G (AGC ÆŢ SE

GGCGTA AATTCAGGTTATGGC TYR AAC TYR ASIN 1460 GCCATTGGGGCA

GAT (

ASP

CTA 日 AAAGACCGC 1520 ARG ASP CCAA GIN AAACA GAA

FIG. 10F

GLU TYR VAL TYR ASP SER LYS GLY GLU ASN GAATATGTTTATGACAGCAAAGGTGAAAT 1540 1550 1560	LYS TRP PHE ASP ASP VAL ARG VAL SER TYR A A T G G T G T G T G T C T T A T 1590 ASP LYS GIN ASP ILE THR LEU ARG SER GIM G A C A A G C A A G A C A T T A C G T A G C C A G 1600 1600	LEU THR ASN THR HIS CYS SER THR TYR PRO CTGACCAACACGCACTGTTCAACCTATCCG 1650 1630 HIS ILLE ASP LYS ASN CYS THR PRO ASP VAL CACATTGACAAAATTGTACGCCTGATGTC 1680	ASN LYS PRO PHE SER VAL LYS GLU VAL ASP A A T A A A C C T T T T C G G T A A A G A G G T G A T 1690 1700 ASN ASN ALA TYR LYS GLU GLN HIS ASN LEU A A C A A T G C C T A C A A G C A C A A T T T A 1740
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FIG. 10

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	T C A C 1790		ARG C G T
5	HLS CAT		TYR I A T
UTC	C A T		ASP 3 A T
THT.	A C G (GLU ; A A (
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LYS 1 A A 1 LEU	r T G (ASN A T T	LEU TTA
N LYS CAAA1		PHE TCA 20	
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FIG. 10K

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ASN A A T 30		ASP G A T		ASP G A T 0		PRO
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FIG. 10F

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GLU GLN GLU HIS ALA LYS ILE ASN THR A A A C A G A A G A A C A T G C C A A A A T C A A T A C A A 1080 1060		LYS HIS ASN PRO PHE ASP ASN SER ILE TRP G A G C A T A A T C C A T T T G A C A A C T C T A T T T G G C 1120 1140		THR VAL TYR ASN GIN GLU LYS GIN ASN ILE G CTGTTTACAACCAAGAAACAATG		ARG PRO ASP LYS LEU ASP ASP VAL ALA L G C C T G A C A A A A A C T T G A T G A C G T G G C A C 1240 1250
ALA G C C		ASP G A C		GLU GAG		LEU CTT
HIS CAT		PHE T T T		GIN C A A		LYS A A A
GLU G A A 1060	. A 10	PRO C C A 1120	A 70	ASN A A C 180	ر 20	LYS A A A 240
GLU GAA	LEU TTG	ASN A A T	GIN CAAA	TYR TAC	GIN C A A 12:	ASP G A C
GLN C A G	ASP	HIS C A T	LYS GLU VAL GIN AAAGAAGTACA 1160	VAL G T T	GLU ASN LYS GIN GAAAATAAACA 1220	PRO C C T
GLU A A	GLY G G T	LYS A G	CLU GAA	THY C T	ASN A A T	ARG G C
	GLU G A A 1100		LYS A A A 1160		GLU G A A 1220	
	LEU CTT		SER AGC		ARG A G A	
	LYS A A A		LYS ASN SER AAAATAG SO		LYS A A A	
	VAL G T A 1090		LYS A A A 150		ILE A T C 210	
	VAL GTT		ILE ATC 1		GLN C A A	
	SN ASP VAL VAL LYS LEU GLU GLY ASP LEU A T G A T G T A A A A C T T G A A G G T G A C T T G A I 1000 1110		IN ASN ILE LYS ASN SER LYS GLU VAL GIN A A A A C A T C A A A A A T A G C A A A A A G A A G T A C A A A A A A A A A A A A A A A A A		LU ASP GIN IIJE LYS ARG GLU ASN LYS GIN A A G A T C A A A A A G A G A A A A A A C A A C 1210 1220	
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FIG. 116

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LYS A A A		LYS A A A		SER A G T		ASN
ALA G C T		ALA G C C		TYR T A T		ASP
THR A C T 1490		ASP G A T 1550		ARG C G T 1610		PRO
GLN C A A		THR ACC		ASP GAT		LYS
THR ACA		MET A T G		GLY GGT		ASIN
PHE TYR GIN GLY THR GIN THR ALA LYS GIN L TTTATCAAGGTACACAAACTGCTAAACAAT 1480 1490 1500	EU PRO VAL SER GIN VAL LYS TYR LYS GLY TGCCTGTATCTCAAGTTAAGTATAAAGGCA 1510 1520	THR TRP ASP PHE MET THR ASP ALA LYS LYS GCTTGGGATTTTATGACCGATGCCAAAAAG	LY GIN SER PHE SER SER PHE GLY THR SER GACAATCATTTAGCAGTTTTGGTACATCGC 1570 1580 1590	GIN ARG LEU ALA GLY ASP ARG TYR SER ALA M A A C G T C T T G C T G G T G T T A T A G T G C A A 1600 1610 1620	ET SER TYR HIS GLU TYR PRO SER LEU LEU TGTCTTACCATGAATACCCATCTTATTAA 1630 1640	THR ASP GLU LYS
	EU PROTECT		LY GLA GACA		ET SER TGTC	

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FIG.11H

AAGAGC AAA GATTTT AGTGAG ີ ປ AGC HIS GAATAT

C G C TAAA AAAACC AAT GTT VAL GGCAGT GLY

ASN ILE TYR GLY ASN AATATCTACGGCAACC 1820

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SUBSTITUTE SHEET (RULE 26)

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PHE THR SER ASP ALA LYS ASN SER LEU GLU G TTACCAGCGATGCCAAAATAGCCTAGAAG 1900 1910 1920	LY GLY PHE TYR GLY PRO ASN ALA GLU GLU GCGGTTTTTAGGACCAAGGAGC 1930 1940 1950	LEU ALA GLY LYS PHE LEU THR ASN ASP ASN L TGGCAGGTAAATTCCTAACCAATGACAACA	YS LEU PHE GLY VAL PHE GLY ALA LYS ARG A A C T C T T T G G T G C T A A A C G A G 1990 2000 2000	GLU SER G AGAGTG	LE LEU ASP ALA TYR ALA LEU GLY THR PHE TCTTAGATGCACTTGGGACATTTA 2050 2060 2060	ASN LYS PRO GLY THR THR ASN PRO ALA PHE TA TA A A C C T G G T A C G A C C A A T C C G C C T T T A 2080 2090 2100

FIG. 11.

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SER AGTA	PRO C C A	THR ACA	ASP PHE G 3 A C T T T G 2520
LEUCTT	VAL G T A	ASP G A C	ILE A T T (
GLU G A G (2330	ALA G C C 2390	LYS A A G (2450	ASP G A C 2510
GLU TYR LEU LYS PHE GLY GLU LEU SER ILE CAATACTTGGTGAGCTTAGTATCG 2330 2340	LEU GIN GLY TACAAGGCG 2370 LU ARG THR ALA GLU LYS ALA VAL PRO THR GAAAGCCGTACCAACCG 2380 2380 2400	GLY ASN TRP 3 G G A A C T G G G 2430 VAL GLY TYR ILE THR GLY LYS ASP THR GLY T T A G G A T A C A T C A C A G G A A A G G A C A C A G G A A 2440 2440 2460	ASN GLU ALA A A T G A G C C C 2490 GIN ASP ILE ALA ASP PHE ASP ILE ASP PHE (A A G A T A T T G C T G A T T T T G A C A T T T G A C A T T T G A C A T T G A C T T T G A C A T T G A C T T T G A C A T T T G A C A T T T G A C A T T T G A C A T T T G A C A T T T G A C A T T T G A C T T T T G A C T T T T T T T T T T T T T T T T T T
PHE TTT	GLU GAG	THR ACA	ASP G A T
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	PHE T T T 2360	LEU C T G 2420	PHE T T T 2480
	VAL G T C	TYR T A T	SER A G C
	SER A G C	LYS A A A	LYS A A A
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10 20 30 MNQSKQNLKSKKSKQVLKLSALSIGILINITQX .Q.QHLFRNILCM .Q.QHLFRNILC			110 120 130 GIAVVEQGRGASSGYSIRGMDKNRVAVLVDGINQ	SLT. VS. SLT. LA. SLT. SLT. SLT. SLT. SLT. SLT. SLT. SLT

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470 480 490 500 STYPHIDKNCTPDVNKPFSVKEVDNNAYKEQHNLJKAVFNRGY.FYKS.RMI.E.SRFQK .ADGSY.R.SADYYKS.RVI.G.S.R.LQ.A.K .ADGSY.R.SADYYKS.RVI.G.S.R.LQ.A.K .LNPSR.TLD.Y.YYKS.R.VKMLQINI.E	510 520 530 540 550 560 THHHINLOVGYDKENSSLSREDYRLATHQSYQKLDYTPPSNPLPDKF-KPILGSNN	TR. NLSINL R. K. Q HS Y. QWAVQAYD. I KP. F. NGS D UR. NKSVNK. F. R. S. B. RHQ YYQHAWRAYSSK KTAN. NGD S TR. NLSVNL T. G. N. RHQ YYQSAWRAYS. K Q. NGKTS PN. REK LT. Q. VFNL. F. D. T. A. QHK TRRVIATA SI . RK GETG . RN. LQS 570

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OKTINIDKIDYQALIDQYDKQNPNSTLKPFEKIKQSLGQEKYNKIDELGFKAYKDLRNEMAGMT	670 680 690 700 NINSQQNANKGRIDNI YQPNQA-TVVKIDIKCKYSETINS-Y	T.NTSPI.RFGNTGN.TGQI.LFGNT	710 720 730 740 750 760 ADCSTTRHISGDNYFIALKINMTINKYVDLGLGARYDRIKHKSDVPLVINSASNQLSMNFGVV	TP.N.G.NG.YA.VQ. VRLGRWA.V.A.IYRSTH.EDKS.STGTHRNA. TP.S.N.KS.YA.VR.VRLGRWA.V.A.LYRSTH. DGS.STGTHRTA.I. TP.S.N.KS.YA.VR.VRLGRWA.V.A.LYRSTH.DGS.STGTHRTA.I. RKV.L.K.K.YF.ARN.ALGI.VSRT.ANESTISVGKFKNFT.I.

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4223 Q8 B16B6 M982 FA19 Eagan		4223 Q8 B16B6 M982 FA19 Eagan
770 780 790 800 VKPTIWILDIAYRSSQSFRMPSFSEMYGERFGVTIGKG FT.M.LT. A.T. L. A. W.A. ESLKTL L. AD. LT. T.T. L. A. W.S. OSKAV L. AD. LT. T.T. L. A. W.S. DK. KAV I. E. LS. L.T. N. WY. GKNDEV	940 EIGATTHNHLGSLET A.A. IVFKGDF.N A. IVFKGDF.N A. IVFKGDF.N A. IVFKGDF.N	### ### ### ### ### ### ### ### ### ##

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910 920 930 940 950 960 INAVNSRLPYGLYSTLAYNKVDVKGKTTINPTLAG-TNILFDALQPSRYVVGLGYDAPSQKWGA WHG.WGG.DRIKDADIRADRTFV.SYV.LH.DGII WNG.WDK.E.WFR.H.RDIKKRADRTDIQSHSQ.EGV F.GLWK.IW.A.FQ.KDQKI.AGSVSSYIIH.NTI	970 980 990 1000 NAIFTHSDAKNPSELLADKNLGNGNIQ-TKQATKAKSTP	TM. Y.K. SVDGSQA.LANAK.A-ASRRTR. GML.Y.K. EITGSRA.LSRNA ARRTR. GML.Y.K. EITGSRA.LSRNA ARRTR. TMQ.KSQNGKRASRDV.S RKLIRA	1010 1020 1030 1040 1050 1060 1070 MQTLDLSGYVNIKDNFTILRAGVYNVFNIYYTTIWEALRQTAEGAVNQHTGLSQDKHYGRYAAFGRNYQLALEMKF* 4223	YVT.V. Y. KHL LL.YR.V. NV. G KNVGV.N. TFS. * B16B6 YIV.V. YTV.KH. LL.YR.V. NV. G KNVGV.N. TFS. * FA19 YIV.V. YTV.KH. LL.HR.V. NV. A KNVGV.N. TFS. * FA19 HIV. YMANK. IM. L. L. YR.V. V. Q QNVGS.T. S T.T. * Eagan

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Tbp2 comparison

10 20 30 40 50 60 MKHIPLTILCVALSAV-LLTACGGS-GCSNPPAPTPI NALSGSGNIGNTGNGGTINT-ANAG	GYGMALSKINI HINRODTPLD-EKNI ITLDGKKQVAEG-KKSPLPFS-LDV-ENKLLDGY IA VELRIMI P. EQEEH-A. INNVV. LEGDL HN. FDN. IWONIK. SKEVQTVY F.VLPRR. AHFN. KYKHKP. GSM. WLQRGEPNSFS. RDE. E F.M-RLKRR. WYPGAE. SEVK. NES. WEATGLPTKP. E KRQKS. I. KVET D-S F.M-RFKRR. WHPSANPK. DEVK. KND. WEATGLPTEP. K LKQQS. ISEVETN. N-S F.M-RFKRR. WHPSANPK. DEVK. KND. WEATGLPTEP. K LKQQS. ISEVETN. N-S
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4223 Q8 B16B6 M982 FA19 Eagan	4223 Q8 B16B6 M982 FA19 :
160 170 180 190 200 KMNADKNAIGDRIKKGNKEISDEELAKQIKEAVRKSHEFQQV- NQEKQNIEDQIK. EN. QRPDKKLDDV. L. AYIEKVLDDRLITELA	210 220 250 LSSLENKIFHSNDGTTKATTRDLKYVDYGY-Y1ANDANYLIVKTDKIMNLGPVGGVFYNGTTT KPIY. KN. NY. H. KQN R RS I. RSGYSITPK. IAKT. FD. AL Q Q

FIG. 130

	4223 Q8 B16B6 M982 FA19 Eagan	
310 320 340 360 360 360 360 360 360 360 360 360 36	VIKTERYDIDANIHGNRFRGSATASNKNDTSK-HPFTSDAN N.KYDITEASKK QITT.Q.TLK.K.L.ADGA.NGSI.SD KHT.QY.SL.Q.TN.TTD.K-ENET.LV.SS YY.SL.TLRS.K.I.TD.PNTGGT.LVF.SS KK.LD.YSTVKPTESEEEGT	410 420 430 NRLEGGFYGPKGEFLAGKFLTNDNFGFGAKRESKAEEKTES. NA EK EKS VAA QKD. KDGENA. GPA SSF. Q GFRSD. Q. VAV. GSTKD. LENGAA. SGS. G-AAASGGAAGTISSE SSF. Q GFRSD. G. VAV. GSTKD. ST NANAP-AASSGFGAATMPS NAG ATRV SETEETKKEALSK. TLIDGKLITFSTKKTDA

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4223 Q8 B16B6 M982 FA19		4223 Q8 B16B6 M982 FA19 Eagan
450 460 470 480AILDAYALGTFNTSNATTFTPFTFKQLDNFGNAKKLVTVIRITGEEFKKE.I.S.DVL NSKLTTVVE.TLNDKKI.NS.AQ ETRLTTVVE.TPDGKEI.NS.TR KTNATTSTA.NTTTDTTANTI.DEKN.KTEDISSE.DY.L.	490 IGSTVIDLVPTDATKNEFTKDKPESAINEAGETLMNDEVSV VDGVELS. LSE-GNKAA	

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	4223 Q8 B16B6 M982 FA19 Eagan	
550 570 SVFLQGERTATTGEKAVPTTGTAKYLG SAMQAGGNSSQADAKTEQVEQ.MVPVSDVAR.EANR. SVMQAVKNSSQADAKTTEQVEQ.MD.RI.QEQGIV KETETETETETETETEXDKEKEKEKDKDKEKQTAATTNTYYQL.HPKDDI.K.SH.	NWGYIT-GKDIGIGKSFTDAQDVADFTIDFGNKSVSGK	620 630 640 650 670 LITKGRQDPVFSITGQLAGNGWIGTASTTKADAGGYKIDSSSTGKSLAIKDANVTGGFYG .T.QNANVVEN.KTA.D.TS.A.TAM.KDFS.V.KGEN.FAL.PQN.N.HYTHE.T.STAEN.AQT.T.E.M.QFEKAES.FDL.QKN.TRTPKAY.T.K.KTAEN.SEAT.T.DAM.EFKKGND.FAP.QNNSTVTHKVH.AN.E.QKRHDTGNEANFNNSS.AFTANFV.GKNSQNKNTPINITTK.N.A.

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4223 Q8-B16B6 M982-FA19 Eagan

PNANEMGGSFT-----NADDSKASV

K. I.SFPGNAPEGKQE-----HDT.

K. E. L. W. AYPGDKQTEKATATSSDG--. SAS. -. T.

E. L. W. AYPGNAPTNSESSSTVSSGS GKNAP A

K. S. L. YNGNSTATNSESSSTVSSGS GKNAP A

-1G.13F

700 VFGTKRQQEV-K*

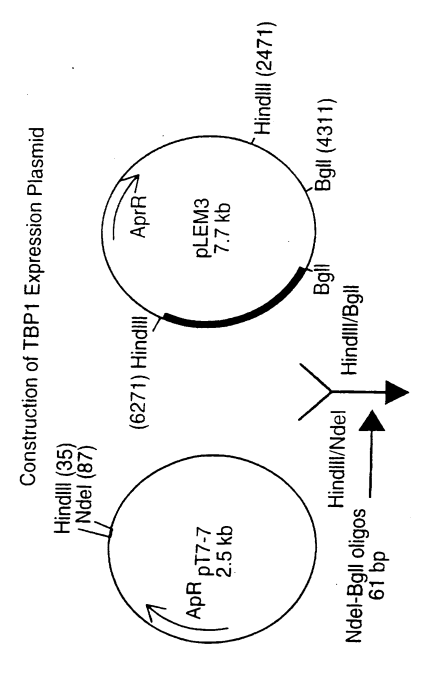
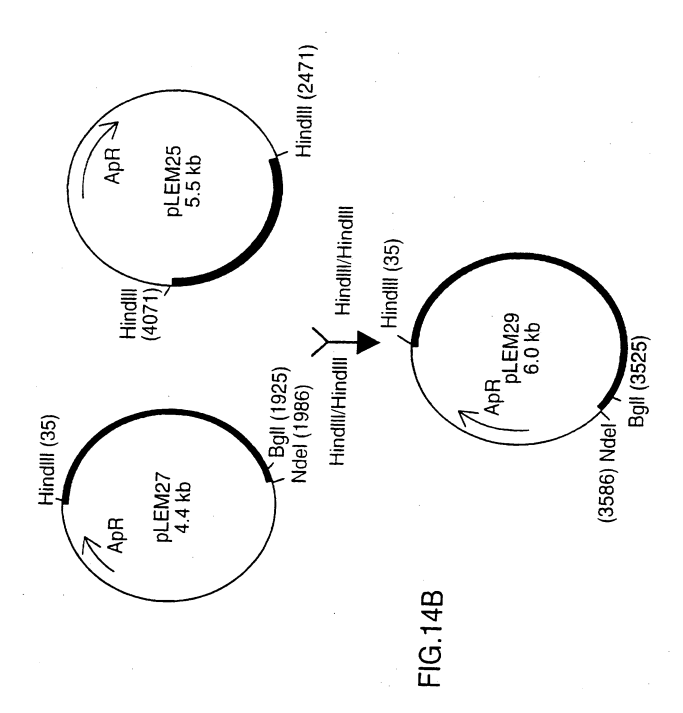
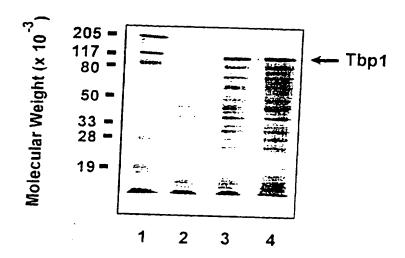


FIG.14A



Expression of rTbp1 in E. coli



- 1. Prestained molecular weight markers
- 2. pLEM29B-1 lysate, non-induced
- 3. pLEM29B-1 lysate, 1 hr post-induction
- 4. pLEM29B-1 lysate, 3 hr post-induction

Fig.15

Purification of Tbp1 from E.Cole

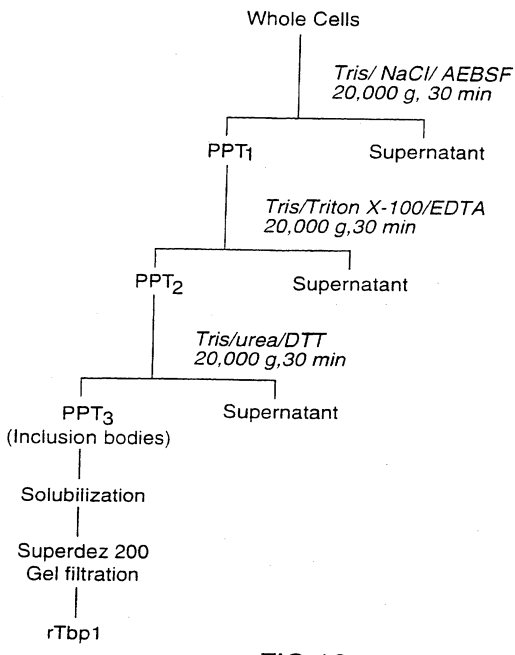
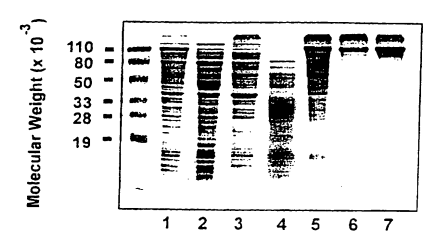


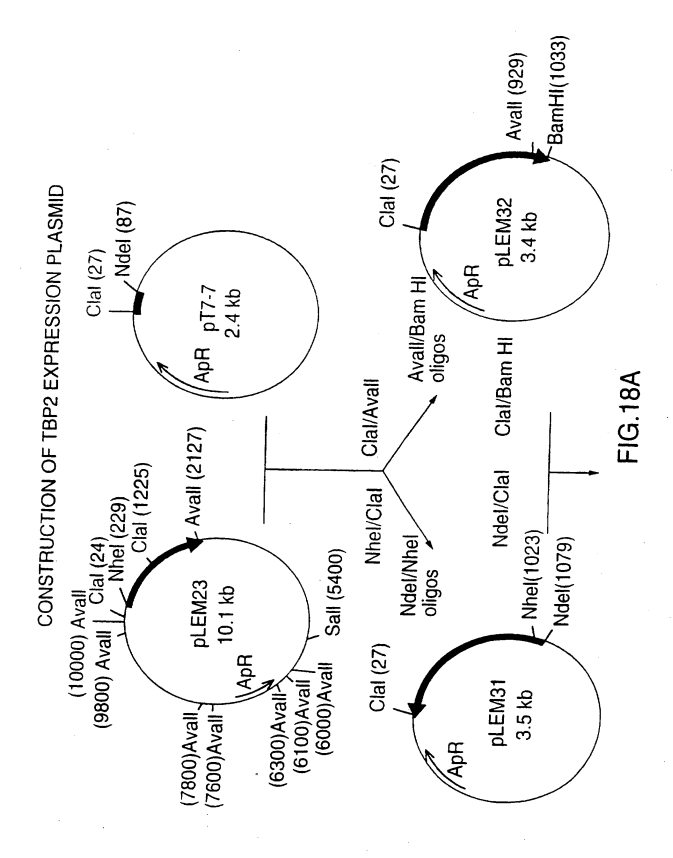
FIG.16

Purification of rTbp1 from E. coli

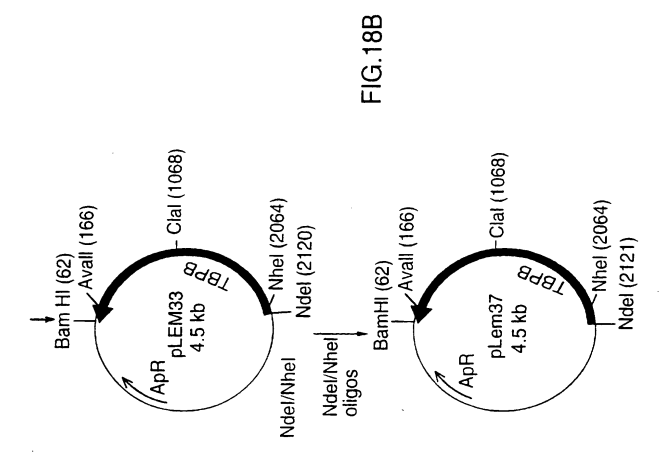


- 1. E. coli Whole cells
- 2. Soluble proteins after 50 mM Tris/ NaCl extraction
- 3. Soluble proteins after Tris/ Triton X-100/ EDTA extraction
- 4. Soluble proteins after Tris/ urea/ DTT extraction
- 5. Left-over pellet (rTbp1 inclusion bodies)
- 6.7. Purified rTbp1

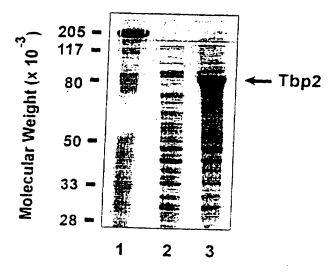
Fig.17



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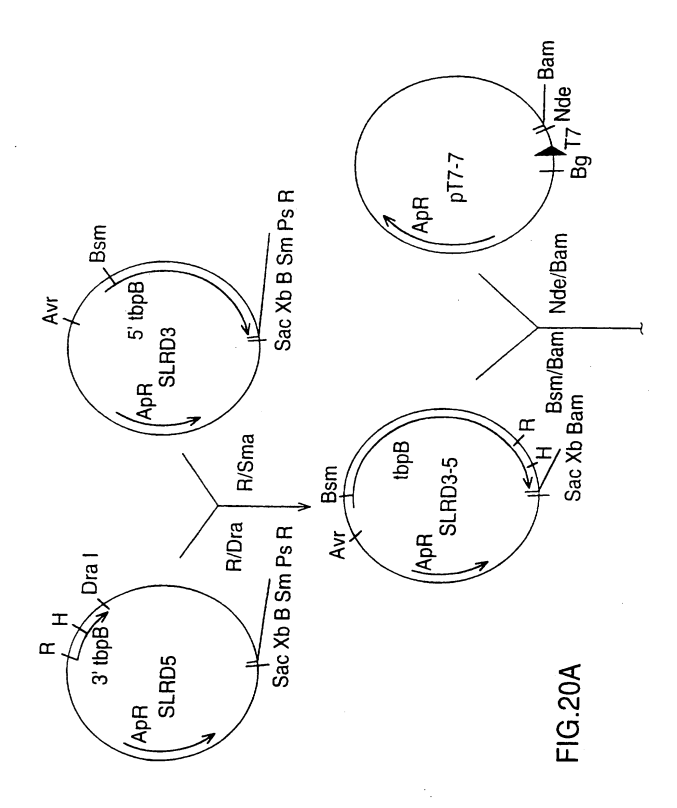


Expression of rTbp2 in E. coli

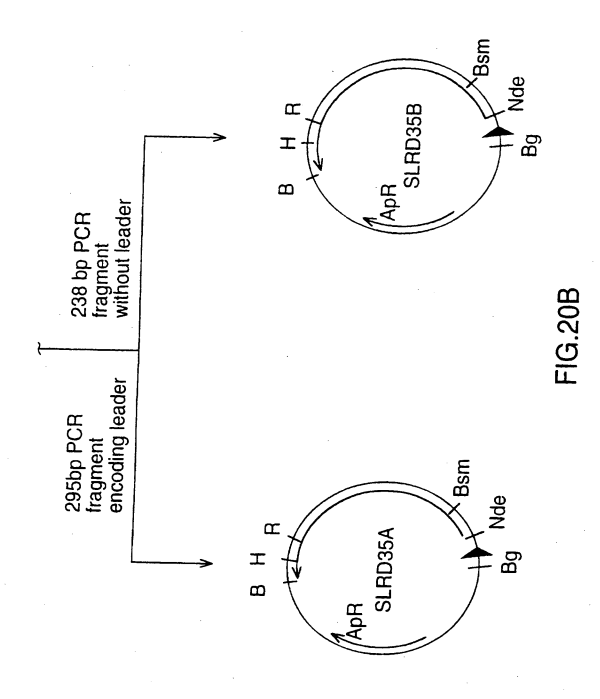


- 1. Prestained molecular weight markers
- 2. pLEM37B-2 lysate, non-induced
- 3. pLEM37B-2 lysate, induced

Fig.19

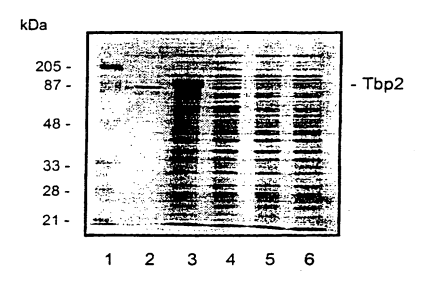


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Fig 21. Expression of Q8 rTbp2 protein in E. coli



- 1. Prestained molecular weight markers
- 2. 4223 rTbp2 protein
- 3. SLRD35A lysate, 3 hr post-induction
- 4. SLRD35B lysate, 3 hr post-induction
- 5. SLRD35A lysate, non-induced
- 6. SLRD35B lysate, non-induced

Purification of Tbp2 from E.Coli

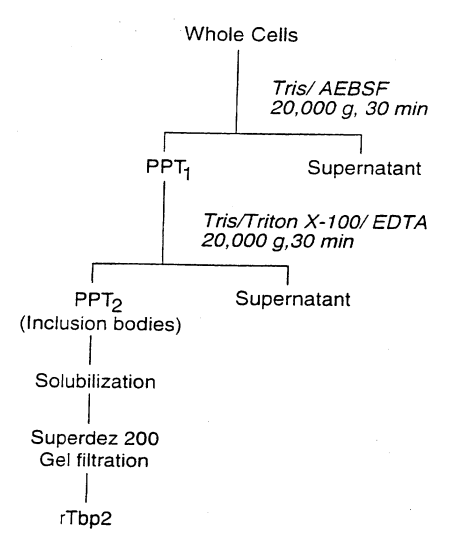
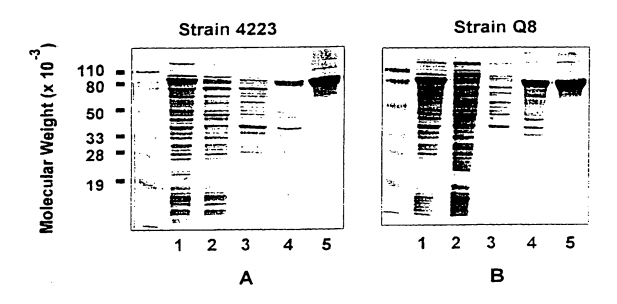


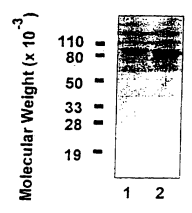
FIG.22

Purification of rTbp2 from E. coli



- 1. E. coli Whole cells
- 2. Soluble proteins after 50 mM Tris extraction
- 3. Soluble proteins after Tris/ Triton X-100/ EDTA extraction
- 4. Left-over pellet (rTbp2 inclusion bodies)
- 5. Purified rTbp2

Binding of Tbp2 to Human Transferrin



- 1. rTbp2 (strain 4223)
- 2. rTbp2 (strain Q8)

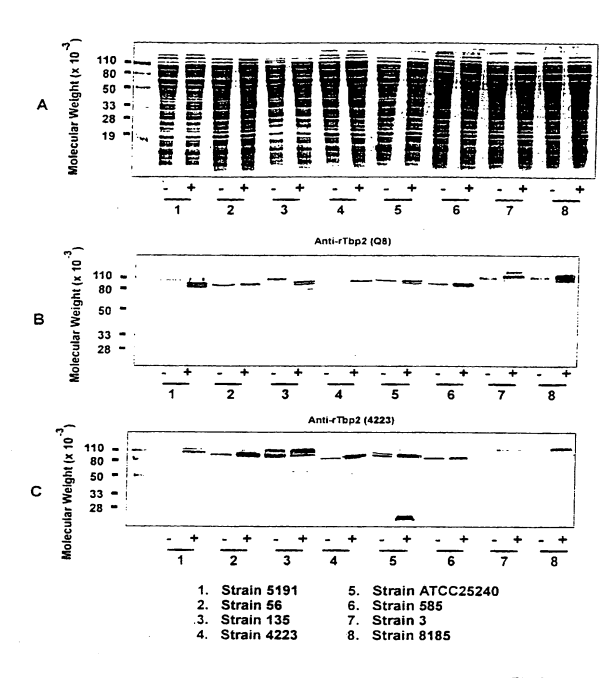


Fig.25





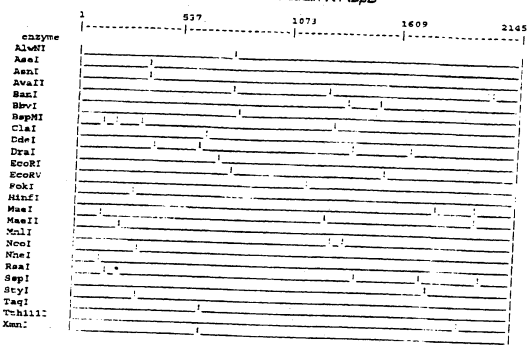


Figure 77 Nucleotide and deduced amino acid sequence of M. catarrhalis R1 tbpB TGTCAGCATGCCAAAATAGGCATTAACAGACCTTTTTTAGATAACACTCATCAGCCCATCAGAGGATTATTTT ATG AAA CAC ATT CCT TTA ACC ACA CTG TGT GTG GCA ATC TCT GCC GTC TTA TTA MET Lys His Ile Pro Leu Thr Thr Leu Cys Val Ala Ile Ser Ala Val Leu Leu ACC GCT TGT GGT GGC AGT GGT TCA AAT CCA CCT GCT CCT ACG CCC ATT CCA Thr Ala Cys Gly Gly Ser Gly Gly Ser Asn Pro Pro Ala Pro Thr Pro Ile Pro AAT GCT AGC GGT TCA GGT AAT ACT GGC AAC ACT GGT AAT GCT GGC GGT ACT GAT Asn Ala Ser Gly Ser Gly Asn Thr Gly Asn Thr Gly Asn Ala Gly Gly Thr Asp 216 AAT ACA GCC AAT GCA GGT AAT ACA GGC GGT ACA AGC TCT GGT ACA GGC AGT GCC Asn Thr Ala Asn Ala Gly Asn Thr Gly Gly Thr Ser Ser Gly Thr Gly Ser Ala 243 AGC ACG TCA GAA CCA AAA TAT CAA GAT GTG CCA ACA ACG CCC AAT AAC AAA GAA Ser Thr Ser Glu Pro Lys Tyr Gln Asp Val Pro Thr Thr Pro Asn Asn Lys Glu 297 324 CAA GIT TCA TCC ATT CAA GAA CCT GCC ATG GGT TAI GGC ATG GCT TTG AGT AAA Gln Val Ser Ser Ile Gln Glu Pro Ala MET Gly Tyr Gly MET Ala Leu Ser Lys 351 ATT AAT CTA TAC GAC CAA CAA GAC ACG CCA TTA GAT GCA AAA AAT ATC ATT ACC Ile Asn Leu Tyr Asp Gln Gln Asp Thr Pro Leu Asp Ala Lys Asn Ile Ile Thr 405 432 TTA GAC GGT AAA AAA CAA GTT GCT GAC AAT CAA AAA TCA CCA TTG CCA TTT TCG Leu Asp Gly Lys Lys Gln Val Ala Asp Asn Gln Lys Ser Pro Leu Pro Phe Ser 459 TTA GAT GTA GAA AAT AAA TTG CTT GAT GGC TAT ATA GCA AAA ATG AAT GAA GCG Leu Asp Val Glu Asn Lys Leu Leu Asp Gly Tyr Ile Ala Lys MET Asn Glu Ala 513 GAT AAA AAT GCC ATT GGT GAA AGA ATT AAG AGA GAA AAT GAA CAA AAT AAA AAA Asp Lys Asn Ala Ile Gly Glu Arg Ile Lys Arg Glu Asn Glu Gln Asn Lys Lys ATA TCC GAT GAA GAA CTT GCC AAA AAA ATC AAA GAA AAT GTG CGT AAA AGC CCT Ile Ser Asp Glu Glu Leu Ala Lys Lys Ile Lys Glu Asn Val Arg Lys Ser Pro

GAG TTT CAG CAA GTA TTA TCA TCG ATA AAA GCG AAA ACT TTC CAT TCA AAT GAC Glu Phe Gln Gln Val Leu Ser Ser Ile Lys Ala Lys Thr Phe His Ser Asn Asp



ANA DCA 100 and 100
AAA ACA ACC AAA GCA ACC ACA CGA GAT TTA AAA TAT GTT GAT TAT GGT TAC TAC Lys Thr Thr Lys Als Thr Thr Arg Asp Leu Lys Tyr Val Asp Tyr Gly Tyr Tyr
729 TIG GTG AAT GAT GCC AAT TAT CTA ACC GTC AAA ACA GAC AAC CCA AAA CTT TGG Leu Val Asn Asp Ala Asn Tyr Leu Thr Val Lys Thr Asp Asn Pro Lys Leu Trp
AAT TCA GGT CCT GTG GGC GGT GTG TIT TAT AAT GGC TCA ACG ACC GCC AAA GAG ASN Ser Gly Pro Val Gly Val Phe Tyr Asn Gly Ser Thr Thr Ala Lys Glu
837 CIG CCC ACA CAA GAT GCG GTC AAA TAT AAA GGA CAT TGG GAC TTT ATG ACC GAT Leu Pro Thr Gln Asp Ala Val Lys Tyr Lys Gly His Trp Asp Phe MET Thr Asp
918 GTT GCC AAA AAA AGA AAC CGA TTT AGC GAA GTA AAA GAA ACC TAT CAA GCA GGC Val Ala Lys Lys Arg Asn Arg Phe Ser Glu Val Lys Glu Thr Tyr Gln Ala Gly
945 TGG TGG TAT GGG GCA TCT TCA AAA GAT GAA TAC AAC CGC TTA TTA ACC AAA GCA Trp Trp Tyr Gly Ala Ser Ser Lys Asp Glu Tyr Asn Arg Leu Leu Thr Lys Ala
999 1026 GAT GCC GCA CCT GAT AAT TAT AGC GGT GAA TAT GGT CAT AGC AGT GAA TTT ACT Asp Ala Ala Pro Asp Asn Tyr Ser Gly Glu Tyr Gly His Ser Ser Glu Fhe Thr
1080 GTT AAT TTT AAG GAA AAA AAA TTA ACA GGT GAG CTG TTT AGT AAC CTA CAA GAC Val Asn Phe Lys Glu Lys Lys Leu Thr Gly Glu Leu Phe Ser Asn Leu Gln Asp
AGC CAT AAA CAA AAA GTA ACC AAA ACA AAA CGC TAT GAT ATT AAG GCT GAT ATC Ser His Lys Gln Lys Val Thr Lys Thr Lys Arg Tyr Asp Ile Lys Ala Asp Ile
1161 CAC GGC AAC CGC TTC CGT GGC AGT GCC ACC GCA AGC GAT AAG GCA GAA GAC AGC His Gly Asn Arg Phe Arg Gly Ser Ala Thr Ala Ser Asp Lys Ala Glu Asp Ser
1215 AAA AGC AAA CAC CCC TTT ACC AGC GAT GCC AAA GAT AAG CTA GAA GGT GGT TTT Lys Ser Lys His Pro Phe Thr Ser Asp Ala Lys Asp Lys Leu Glu Gly Gly Phe
1269 TAT GGA CCA AAA GGC GAG GAG CTG GCA GGT AAA TTC TTA ACC GAT GAT AAC AAA Tyr Gly Pro Lys Gly Glu Glu Leu Ala Gly Lys Phe Leu Thr Asp Asp Asn Lys
1323 CTC TTT GGT GTC TTT GGT GCC AAA CAA GAG GGT AAT GTA GAA AAA ACC GAA GCC Leu Phe Gly Val Phe Gly Ala Lys Gln Glu Gly Asn Val Glu Lys Thr Giv Ala

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1377

ATC TTA GAT GCT TAT GCA CIT GGG ACA TIT AAT AAA CCT GGT ACG ACC AAT CCC Ile Leu Asp Ala Tyr Ala Leu Gly Thr Phe Asn Lys Pro Gly Thr Thr Asn Pro

1431 1458

GCC TTT ACC GCT AAC AGC AAA AAA GAA CTG GAT AAC TTT GGC AAT GCC AAA AAG Ala Phe Thr Ala Asn Ser Lys Lys Glu Leu Asp Asn Phe Gly Asn Ala Lys Lys

1485 2512

TTG GTC TTG GGT TCT ACC GTC ATT GAT TTG GTG CCT ACT GAT GCC ACC AAA GAT Leu Val Leu Gly Ser Thr Val Ile Asp Leu Val Pro Thr Asp Ala Thr Lys Asp

1539

GTC AAT GAA TTC AAA GAA AAG CCA AAG TCT GCC ACA AAC AAA GCG GGC GAA ACT Val Asn Glu Phe Lys Glu Lys Pro Lys Ser Ala Thr Asn Lys Ala Gly Glu Thr

1593

TTG ATG GTG AAT GAT GAA GTT AGC GTC AAA ACC TAT GGC AAA AAC TTT GAA TAC Leu MET Val Asn Asp Glu Val Ser Val Lys Thr Tyr Gly Lys Asn Phe Glu Tyr

1647

CTA AAA TIT GGT GAG CIT AGT GTC GGT AGC CAT AGC GTC TIT TTA CAA GGC Leu Lys Phe Gly Glu Leu Ser Val Gly Gly Ser His Ser Val Phe Leu Gln Gly

1701 1728

GAA CGC ACC GCT ACC ACA GGC GAG AAA GCC GTA CCA ACC ACA GGC AAA GCC AAA GLU Arg Thr Ala Thr Thr Gly Glu Lys Ala Val Pro Thr Thr Gly Lys Ala Lys

1755

TAT TTG GGG AAC TGG GTA GGA TAT ATC ACA GGA GCG GAC TCA TCA AAA GGC TCT Tyr Leu Gly Asn Trp Val Gly Tyr Ile Thr Gly Ala Asp Ser Ser Lys Gly Ser

1809

ACC GAT GGC AAA GGC TIT ACC GAT GCC AAA GAT ATT GCT GAT TIT GAC ATT GAC Thr Asp Gly Lys Gly Phe Thr Asp Ala Lys Asp Ile Ala Asp Phe Asp Ile Asp

1863

TTT GAG AAA AAA TCA GTT AAT GGC AAA CTG ACC ACC AAA GAC CGC CAA GAC CCT Phe Glu Lys Lys Ser Val Asn Gly Lys Leu Thr Thr Lys Asp Arg Gln Asp Pro

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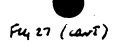
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ACC GCC GAA GCG AAC GCA GGG GGC TAT AAG ATA GAT TCT AGC AGT ACA GGC AAA Thr Ala Glu Ala Asn Ala Gly Gly Tyr Lys Ilê Asp Ser Ser Ser Thr Gly Lys

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TCC ATC GTC ATC AAA GAT GCC GTG GTT ACA GGT GGC TTT TAT GGT CCA AAT GCA Ser Ile Val Ile Lys Asp Ala Val Val Thr Gly Gly Phe Tyr Gly Fro Asn Ala



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ACC GAG ATG GGT GGG TCA TIT ACA CAC AAC AGC GGT AAT GAT GGT AAA STC TCT Thr Glu MET Gly Gly Ser Phe Thr His Asn Ser Gly Asn Asp Gly Lys Val Ser

2133

GTG GTC TTT GGC ACA AAA AAA CAA GAA GTT AAG AAG TGA Val Val Phe Gly Tor Lys Lys Gln Glu Val Lys Lys *

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DFGNKSVSGKLITKGRQDPVFSITGQIAGNGWTGTASTTKADAGGYKIDSSSTGKSIAIKDANVTGGFYGPNANEMGGSFTHNA-DDSKASVVPGTKRQQKVK.*

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In tional Application No PCT/CA 97/00163

G01N33/68

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 6 CO7K C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

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Further documents are listed in the continuation of box C.

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Ε	WO 97 13785 A (CONNAUGHT LAB ; YANG YAN PING (CA); MYERS LISA E (CA); HARKNESS ROB) 17 April 1997 see the whole document	1-25
Y	WO 90 12591 A (UNIV TECHNOLOGIES INTERNATIONA ;SCHRYVERS ANTHONY BERNARD (CA)) 1 November 1990 see claims 1-26	1-25
	-/	

Further documents are listed in the continuation of box C.	X Patent family members are listed in annex.
* Special categories of cited documents:	
'A' document defining the general state of the art which is not considered to be of particular relevance 'E' earlier document but published on or after the international filing date 'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) 'O' document referring to an oral disclosure, use, exhibition or other means 'P' document published prior to the international filing date but later than the priority date claimed	'Y' document of particular relevance; the claimed invention cannot be considered no involve an invention cannot be considered novel or cannot be considered to involve an invention the document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. '&' document member of the same patent family
Date of the actual completion of the international search	Date of mailing of the international search report
17 July 1997	30 JULY 1997 (30.07.97)
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authonzed officer Nauche, S

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E	CA 97/00163	_

		A 37/00103
	non) DOCUMENTS CONSIDERED TO BE RELEVANT	Relevant to claim No.
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	MICROBIAL PATHOGENESIS, vol. 15, 1993, pages 433-445, XP000612196 RAONG-HUA YU ET AL: "THE INTERACTION BETWEEN HUMAN TRANSFERRIN AND TRANSFERRIN BINDING PROTEIN 2 FROM MORAXELLA (BRANHAMELLA) CATARRHALIS DIFFERS FROM THAT OF OTHER HUMAN PATHOGENS" see the whole document	1-25
A	WO 95 33049 A (PASTEUR MERIEUX SERUMS VACC ;TRANSGENE SA (FR); MILLET MARIE JOSE) 7 December 1995 see the whole document	1-25
A	WO 93 08283 A (UNIV SASKATCHEWAN) 29 April 1993 see the whole document	1-25
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nternational application No.

PCT/CA 97/00163

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inc	ernational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. 🗓	Claims Nos.: 23 because they relate to subject matter not required to be searched by this Authority, namely: Remark: Although claim 23 is directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.
2.	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inter	national Searching Authority found multiple inventions in this international application, as follows:
1. A	as all required additional search fees were timely paid by the applicant, this International Search Report covers all earchable claims.
2	is all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment f any additional fee.
3. A	s only some of the required additional search fees were timely paid by the applicant, this International Search Report evers only those claims for which fees were paid, specifically claims Nos.:
4. No	o required additional search fees were timely paid by the applicant. Consequently, this International Search Report is stricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on	The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

on patent family members 97/00163 Publication Patent family Publication Patent document date member(s) date cited in search report AU 7208296 A 30-04-97 WO 9713785 A 17-04-97 WO 9012591 A 01-11-90 US 5292869 A 08-03-94 AU 649950 B 09-06-94 ΑU 5526190 A 16-11-90 EP 03-03-93 0528787 A 26-11-92 JP 4506794 T 24-02-95 NZ 247967 A 25-08-92 US 5141743 A 01-12-95 WO 9533049 A 07-12-95 FR 2720408 A 2675795 A AU 21-12-95 2167936 A 07-12-95 CA EP 0720653 A 10-07-96 FI 960428 A 28-03-96 75992 A 28-05-97 HU JP 9501059 T 04-02-97 NO 960332 A 21-03-96 29-04-93 WO 9308283 A US 5417971 A 23-05-95 AU 2751392 A 21-05-93 2121364 A 29-04-93 CA 17-08-94 EP 0610260 A 5521072 A 28-05-96 US

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